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DNA CHAIN USEFUL FOR XANTHOPHYLL SYNTHESIS AND PROCESS FOR PRODUCING (54)**XANTHOPHYLLS**

The following DNA chains relate to xanthophylls having a keto group, represented by astaxanthin, and the following technique relates to a genetically engineered production of xanthophylls: a DNA chain having a base sequence coding for a polypeptide having an enzymatic activity of converting the 4-methylene group of β -ionone ring into a keto group; a DNA chain having a base sequence coding for a polypeptide having an enzymatic activity of converting the 4-methylene group of a 3-hydroxy-β-ionone ring into a keto group; a DNA chain having a base sequence coding for a polypeptide having an enzymatic activity of adding a hydroxyl group to the 3-carbon atom of a 4-keto- β -ionone ring; and a process for producing various xanthophylls, such a canthaxanthin and astaxanthin, by introducing the above DNA chain(s) into a suitable microorganism, e.g., Escherichia coli, followed by expression thereof.

Description

Technical Field

The present invention relates to DNA strands useful for the synthesis of keto group-containing xanthophylls (keto-carotenoids) such as astaxanthin which are useful for heightening the color of cultured fishes and shellfishes such as sea breams, salmons, lobster and the like and is used for foods as a coloring agent and an antioxidant, and to a process for producing keto group-containing xanthophylls (ketocarotinoids) such as astaxanthin with use of a microorganism into which the DNA strands have been introduced.

Background Art

The term xanthophylls mean carotenoid pigments having an oxygen-containing group such as a hydroxyl group, a keto group or an epoxy group. Carotenoids are synthesized by the isoprenoid biosynthetic process which is used in common halfway with steroids and other terpenoids with mevalonic acid as a starting material. C15 farnesyl pyrophosphate (FPP) resulting from isoprene basic biosynthetic pathway is condensed with C5 isopentenyl pyrophosphate (IPP) to give C20 geranylgeranyl pyrophosphate (GGPP). Two molecules of GGPP are condensed to synthesize a colorless phytoene as an initial carotenoid. The phytoene is converted into phytofluene, ζ -carotene, neurosporene and then lycopene by a series of desaturation reactions, and lycopene is in turn converted into β -carotene by the cyclization reaction. It is believed that a variety of xanthophylls are synthesized by introducing a hydroxyl group or a keto group into the β -carotene (See Britton, G., "Biosynthesis of Carotenoids"; Plant Pigments, Goodwin, T.W. ed., London, Academic Press, 1988, pp. 133-182).

The present inventors have recently made it possible to clone a carotenoid biosynthesis gene cluster from a epiphytic non-photosynthetic bacterium <u>Erwinia uredovora</u> in <u>Escherichia coli</u> with an index of the yellow tone of the bacterium, a variety of combinations of the genes being expressed in microorganisms such as <u>Escherichia coli</u> to produce phytoene, lycopene, β-carotene, and zeaxanthin which is a derivative of β-carotene into which hydroxyl groups have been introduced (See Fig. 10; Misawa, N., Nakagawa, M., Kobayashi, K., Yamano, S., Izawa, Y., Nakamura, K., Harashima, K.; "Elucidation of the <u>Erwinia uredovora</u> Carotenoid biosynthetic Pathway by Functional Analysis of Gene Products Expressed in <u>Escherichia coli</u>, J. Bacteriol., 172, p.6704-6712, 1990; Misawa, N., Yamano, S., Ikenaga, H., "Production of β-carotene in <u>Zymomonas mobilis</u> and <u>Agrobacterium tumefaciencs</u> by Introduction of the Biosynthesis Genes from <u>Erwinia uredovora</u>", Appl. environ. Microbiol., 57, p. 1847-1849, 1991; and Japanese Patent Application No. 58786/1991 (Japanese Patent Application No. 53255/1990): "DNA Strands useful for the Synthesis of Carotenoids").

On the other hand, astaxanthin, a red xanthophyll, is a typical animal carotenoid which occurs particularly in a wide variety of marine animals including red fishes such as a sea bream and a salmon, and crustaceans such as a crab and a lobster. In general, animals cannot biosynthesize carotenoids, so that it is necessary for them to ingest carotenoids synthesized by microorganisms or plants from their environments. Thus, astaxanthin has hitherto been used widely for strengthening the color of cultured fishes and shellfishes such as a sea bream, a salmon, a lobster and the like. Moreover, astaxanthin has attracted attention not only as a coloring matter in foods but also as an anti-oxidant for removing active oxygen generated in bodies, which causes carcinoma (see Takao Matsuno ed., "Physiological Functions and Bioactivities of Carotenoids in Animals", Kagaku to Seibutsu, 28, p. 219-227, 1990). As the sources of astaxanthin, there have been known crustaceans such as a krill in the Antarctic Ocean, cultured products of a yeast Phaffia, cultured products of a green alga Haematococcus, and products obtained by the organic synthetic methods. However, when crustaceans such as a krill in the Antarctic Ocean or the like are used, it requires laborious works and much experfor the isolation of astaxantin from contaminants such as lipids and the like during the harvesting and extraction of the krill. Moreover, in the case of the cultured product of the yeast Phaffia, a great deal of expenses are required for the gathering and extraction of astaxanthin, since the yeast has rigid cell walls and produces astaxanthin only in a low yield. Also, in the case of the cultured product of the green alga Haematococcus, not only a location for collecting sunlight or an investment of a culturing apparatus for supplying an artificial light is required in order to supply light which is essential to the synthesis of astaxantin, but also it is difficult to separate astaxanthin from fatty acid esters as by-products or chlorophylls present in the cultured products. From these reasons, astaxanthin produced from biological sources is in the present situation inferior to that obtained by the organic synthetic methods on the basis of cost. The organic synthetic methods however have a problem of by-products produced during the reactions in consideration of its use as a feed for fishes and shellfishes and an additive to foods, and the products obtained by the organic synthetic methods are opposed to the consumer's preference for natural products. Thus, it has been desired to supply an inexpensive astaxanthin which is safe and produced from biological sources and thus has a good image to consumers and to develop a process for producing the astaxanthin.

Disclosure of the Invention

It would be considered very useful to find a group of genes for playing a role of the biosynthesis of astaxanthin, because it is possible to afford astaxanthin-producing ability to a microorganism optimum in safety as a food or in potentiality for producing astaxanthin, regardless of the presence of astaxanthin-producing ability, by introducing a gene cluster for astaxanthin biosynthesis into the microorganism. No problem of by-products as contaminants is caused in this case, so that it would be considered not so difficult to increase the production amount of astaxanthin with a recent advanced technique of gene manipulation to a level higher than that accomplished by the organic synthetic methods. However, the groups of genes for synthesizing zeaxanthin, one of the xanthophylls, have already been acquired by the present inventors as described above, while no genes encoding a keto group-introducing enzyme required for the synthesis of astaxanthin have not successfully obtained. The reason of the failure in obtaining the genes includes that the keto group-introducing enzyme is a membrane protein and loses its activity when isolated from the membrane, so that it was impossible to purify the enzyme or measure its activity and no information on the enzyme has been obtained. Thus, it has hitherto been impossible to produce astaxanthin in microorganisms by gene manipulation.

The object of the present invention is to provide DNA strands which contain genes required for producing keto group-containing xanthophylls (ketocarotenoids) such as astaxanthin in microorganisms by obtaining such genes coding for enzymes such as a keto group-introducing enzyme required for producing keto group-containing xanthophylls (ketocarotenoids) such as astaxanthin, and to provide a process for producing keto group-containing xanthophylls (ketocarotenoids) such as astaxanthin with the microorganisms into which the DNA strands have been introduced.

The gene cloning method which is often used usually comprising purifying the aimed protein, partially determining the amino acid sequence and obtaining genes by a synthetic probe cannot be employed because of the purification of the astaxanthin synthetic enzyme being impossible as described above. Thus, the present inventors have paid attention to the fact that the cluster of carotenoid synthesis genes in non-photosynthetic bacterium (Erwinia) functions in Escherichia coli, in which lycopene and β-carotene which are believed to be intermediates for biosynthesis of astaxanthin are allowed to produce with combinations of the genes from the gene cluster, and have used Escherichia coli as a host for cloning of astaxanthin synthetic genes. The present inventors have also paid attention to the facts that some marine bacteria have an astaxanthin-producing ability (Yokoyama, A., Izumida, H., Miki, W., "Marine bacteria produced astaxanthin", 10th International Symposium on Carotenoids, Abstract, CL11-3, 1993), that a series of related genes would constitute a cluster in the case of bacteria, and that the gene cluster would be expressed functionally in Escherichia coli in the case of bacteria. The present inventors have thus selected the marine bacteria as the gene sources. They have carried out researches with a combination of these two means and successfully obtained the gene group which is required for the biosynthesis of astaxanthin and the other keto group-containing xanthophylls from marine bacteria. They have thus accomplished the present invention. In addition, it has been first elucidated in the present invention that the astaxanthin synthesis gene cluster in marine bacteria constitutes a cluster and expresses its function in Escherichia coli, and these gene products can utilize β-carotene or lycopene as a substrate.

The DNA strands according to the present invention are set forth as follows.

- (1) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting the methylene group at the 4-position of the β -ionone ring into a keto group.
- (2) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting the methylene group at the 4-position of the β -ionone ring into a keto group and having an amino acid sequence substantially of amino acid Nos. 1 212 which is shown in the SEQ ID NO: 1.
- (3) A DNA strand hybridising the DNA strand described in (2) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (2).
- (4) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting the methylene group at the 4-position of the β -ionone ring into a keto group and having an amino acid sequence substantially of amino acid Nos. 1 242 which is shown in the SEQ ID NO: 5.
- (5) A DNA strand hybridizing the DNA strand described in (4) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (4).
- (6) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting β-carotene into canthaxanthin via echinenone and having an amino acid sequence substantially of amino acid Nos. 1 212 which is shown in the SEQ ID NO: 1.
 - (7) A DNA strand hybridizing the DNA strand described in (6) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (6).
- (8) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting β-carotene into canthaxanthin via echinenone and having an amino acid sequence substantially of amino acid Nos. 1 242 which is shown in the SEQ ID NO: 5.
- (9) A DNA strand hybridizing the DNA strand described in (8) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (8).

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- (10) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting the methylene group at the 4-position of the 3-hydroxy-β-ionone ring into a keto group.
- (11) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting the methylene group at the 4-position of the 3-hydroxy- β -ionone ring into a keto group and having an amino acid sequence substantially of amino acid Nos. 1 212 which is shown in the SEQ ID NO: 1.
- (12) A DNA strand hybridizing the DNA strand described in (11) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (11).
- (13) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting the methylene group at the 4-position of the 3-hydroxy-β-ionone ring into a keto group and having an amino acid sequence substantially of amino acid Nos. 1 242 which is shown in the SEQ ID NO: 5.
- (14) A DNA strand hybridizing the DNA strand described in (13) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (13).
- (15) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting zeaxanthin into astaxanthin by way of 4-ketozeaxanthin and having an amino acid sequence substantially of amino acid Nos. 1 212 which is shown in the SEQ ID NO: 1.
- (16) A DNA strand hybridizing the DNA strand described in (15) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (15).
- (17) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting zeaxanthin into astaxanthin by way of 4-ketozeaxanthin and having an amino acid sequence substantially of amino acid Nos. 1 242 which is shown in the SEQ ID NO: 5.
- (18) A DNA strand hybridizing the DNA strand described in (17) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (17).
- (19) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for adding a hydroxyl group to the 3-carbon of the 4-keto-β-ionone ring.
- (20) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for adding a hydroxyl group to position 3-carbon of the 4-keto- β -ionone ring and having an amino acid sequence substantially of amino acid Nos. 1 162 which is shown in the SEQ ID NO: 2.
- (21) A DNA strand hybridizing the DNA strand described in (20) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (20).
- (22) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for adding a hydroxyl group to position 3-carbon of the 4-keto-β-ionone ring and having an amino acid sequence substantially of amino acid Nos. 1 162 which is shown in the SEQ ID NO: 6.
- (23) A DNA strand hybridizing the DNA strand described in (22) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (22).
- (24) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting canthaxanthin into astaxanthin by way of phoenicoxanthin and having an amino acid sequence substantially of amino acid Nos. 1 162 which is shown in the SEQ ID NO: 2.
- (25) A DNA strand hybridizing the DNA strand described in (24) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (24).
- (26) A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting canthaxanthin into astaxanthin by way of phoenicoxanthin and having an amino acid sequence substantially of amino acid Nos. 1 162 which is shown in the SEQ ID NO: 6.
- (27) A DNA strand hybridizing the DNA strand described in (26) and having a nucleotide sequence which encodes a polypeptide having an enzyme activity described in (26).

The present invention also relates to a process for producing xanthophylls.

That is, the process for producing xanthophylls according to the present invention is set forth below.

- (1) A process for producing a xanthophyll comprising introducing the DNA strand described in any one of the above mentioned DNA strands (1) (9) into a microorganism having a β -carotene-synthesizing ability, culturing the transformed microorganism in a culture medium, and obtaining canthaxanthin or echinenone from the cultured cells.
- (2) A process for producing a xanthophyll comprising introducing the DNA strand described in any one of the above mentioned DNA strands (10) (18) into a microorganism having a zeaxanthin-synthesizing ability, culturing the transformed microorganism in a culture medium, and obtaining astaxanthin or 4-ketozeaxanthin from the cultured cells.
- (3) A process for producing a xanthophyll comprising introducing the DNA strand described in any one of the above mentioned DNA strands (19) (27) into a microorganism having a canthaxanthin-synthesizing ability, culturing the transformed microorganism in a culture medium, and obtaining astaxanthin or phoenicoxanthin from the cultured cells.

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(4) A process for producing a xanthophyll according to any one of the above mentioned processes (1) - (3), wherein the microorganism is a bacterium or yeast.

Brief Description of the Drawings

Fig. 1 illustrates diagrammatically the nucleotide sequence of the keto group-introducing enzyme gene (crt W gene) of the marine bacterium Agrobacterium aurantiacus sp. nov. MK1 and the amino acid sequence of a polypeptide to be encoded thereby.

Fig. 2 illustrates diagrammatically the nucleotide sequence of the hydroxyl group-introducing enzyme gene (crt Z gene) of the marine bacterium Agrobacterium aurantiacus sp. nov. MK1 and the amino acid sequence of a polypeptide

to be encoded thereby.

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Fig. 3 illustrates diagrammatically the nucleotide sequence of the lycopene-cyclizing enzyme gene (crt Y gene) of the marine bacterium Agrobacterium aurantiacus sp. nov. MK1 and the amino acid sequence of a polypeptide to be encoded thereby.

Fig. 4 illustrates diagrammatically the continuation of the sequences following to those illustrated in Fig. 3.

Fig. 5 illustrates diagrammatically the nucleotide sequence of the xanthophyll synthesis gene cluster of the marine bacterium Agrobacterium aurantiacus sp. nov. MK1.

The letters A - F in Fig. 5 correspond to those in Figs. 1 - 4.

Fig. 6 illustrates diagramatically the continuation of the sequence following to that illustrated in Fig. 5.

Fig. 7 illustrates diagrammatically the continuation of the sequence following to that illustrated in Fig. 6.

Fig. 8 illustrates diagrammatically the continuation of the sequence following to that illustrated in Fig. 7.

Fig. 9 illustrates diagrammatically the continuation of the sequence following to that illustrated in Fig. 8.

Fig. 10 illustrates diagrammatically the carotenoid biosynthetic route of the non-photosynthesis bacterium Erwinia uredovora and the functions of the carotenoid synthetic genes.

Fig. 11 illustrates diagrammatically the main xanthophyll biosynthetic routes of marine bacteria Agrobacterium aurantiacus sp. nov. MK1 and Alcaligenes sp. PC-1 and the functions of the xanthophyll synthesis genes.

The function of crtY gene, however, has been confirmed only in the former bacterium.

Fig. 12 illustrates diagrammatically a variety of deletion plasmids containing the xanthophyll synthesis genes (cluster) of the marine bacterium Agrobacterium aurantiacus sp. nov. MK1.

The letter P represents the promoter of the lac of the vector pBluescript II SK. The positions of cutting with restriction enzymes are represented by abbreviations as follows: Sa, Sacl; X, Xbal; B, BamHI; P, Pstl; E, EcoRI; S, Sall; A, Apal; K, Kpnl; St, Stul; N, Nrul; Bg, Bglll; Nc, Ncol; Hc, Hincll.

Fig. 13 illustrates diagrammatically the nucleotide sequence of the keto group-introducing enzyme gene (crtW gene) of the marine bacterium Alcaligenes sp. PC-1 and the amino acid sequence of a polypeptide to be encoded

thereby.

Fig. 14 illustrates diagrammatically the continuation of the sequences following to those illustrated in Fig. 13.

Fig. 15 illustrates diagrammatically the nucleotide sequence of the hydroxyl group-introducing enzyme gene (crtZ gene) of the marine bacterium Alcaligenes sp. PC-1 and the amino acid sequence of a polypeptide to be encoded thereby.

Fig. 16 illustrates diagrammatically the nucleotide sequence of the xanthophyll synthetic gene cluster of the marine bacterium Alcaligenes sp. PC-1 and the amino acid sequence of a polypeptide to be encoded thereby. The letters A -D

in Fig. 16 correspond to those in Figs. 13 - 15.

Fig. 17 illustrates diagrammatically the continuation of the sequences following to those illustrated in Fig. 16.

Fig. 18 illustrates diagrammatically the continuation of the sequences following to those illustrated in Fig. 17.

Fig. 19 illustrates diagrammatically a variety of deletion plasmids containing the xanthophyll synthetic genes (cluster) of the marine bacterium Alcaligenes sp. PC-1.

The letter ${f P}$ represents the promoter of the <u>lac</u> of the vector pBluescript II SK+.

Fig. 20 illustrates diagrammatically xanthophyll biosynthetic routes containing miner biosynthetic routes in the marine bacteria Agrobacterium aurantiacus sp. no. MK1 and Alcaligenes sp. PC-1 and the functions of the xanthophyll synthesis genes.

Miner biosynthetic routes are represented by dotted arrows.

Best Mode for carrying out the Invention

The present invention is intended to provide DNA strands which are useful for synthesizinga keto group-containing xanthophylls (ketocarotenoids) such as astaxanthin derived from a marine bacteria Agrobacterium aurantiacus sp. nov. MK1 and Alcaligenes sp. PC-1, and a process for producing keto group-containing xanthophylls (ketocarotenoids), i.e. astaxanthin, phoenicoxanthin, 4-ketozeaxanthin, canthaxanthin, and echinenone with use of a microorganism into which the DNA strands have been introduced.

The DNA strands according to the present invention are in principle illustrated generally by the aforementioned D** strands (1), (10) and (19) from the standpoint of the fine chemical-generating reaction, and basically defined by the icrementioned DNA strands (2), (4), (11), (13), (20) and (22). The specific examples of the DNA strands (2) and (4) are the aforementioned DNA strands (6) and (8); the specific examples of the DNA strands (11) and (13) are the aforementioned DNA strands (15) and (17); and the specific examples of the DNA strands (20) and (22) are the aforementioned DNA strands (24) and (26). In this connection, the DNA strands (3), (5), (7), (9), (12), (14), (16), (18), (21), (23), (25) and (27) hybridize the DNA strands (2), (4), (6), (8), (11), (13), (15), (17), (20), (22), (24) and (26), respectively, under a stringent condition.

The polypeptides encoded by the DNA strands according to the present invention have amino acid sequences substantially in a specific range as described above in SEQ ID NOS: 1 - 2, and 5 - 6 (Figs. 1 - 2, and 13 - 15), e.g. an amino acid sequence of amino acid Nos. 1 - 212 in SEQ ID NOS: 1 (A - B in Fig. 1). In the present invention, four polypeptides encoded by these DNA strands, that is four enzymes participating in the xanthophyll-producing reaction) may be modified by deletion, substitution or addition in some of the amino acids provided that the polypeptides have the enzyme activities as described above (see Example 13). This corresponds to that "amino acid sequencessubstantially ..." For instance, an enzyme of which amino acid at the first position (Met) has been deleted is also involved in the polypeptide or enzyme obtained by the modification of the amino acid sequence. In this connection, it is needless to say that the DNA strands according to the present invention for encoding the polypeptides also include, in addition to those having nucleotide sequences in a specific range shown in SEQ ID NOS: 1 - 2, and 13 - 15 (Figs. 1 - 2, and 13 - 15), degenerate isomers encoding the same polypeptides as above except degenerate codons.

Keto group-introducing enzyme gene (crtW)

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The DNA strands (1) - (18) are genes which encode the keto group-introducing enzymes (referred to hereinafter as crtW). Typical examples of the genes are crtW genes cloned from the marine bacteria Agrobacterium aurantiacus sp. nov. MK1 or Alcaligenes sp. PC-1, which are the DNA strands comprising the nucleotide sequences encoding the polypeptides having the amino acid sequences A - B in Fig. 1 (amino acid Nos. 1 - 212 in SEQ ID NOS: 1) or A - B in Figs. 13 - 14 (amino acid Nos. 1 - 242 in SEQ ID NOS: 5). The crtW gene product (also referred to hereinafter as CrtW) has an enzyme activity for converting the 4-methylene group of the β-ionone ring into a keto group, and one of the specific examples is an enzyme activity for synthesizing canthaxanthin with β -carotene as a substrate by way of echinenone (see Fig. 11). In addition, the crtW gene product also has an enzyme activity for converting the 4-methylene group of the 3-hydroxy-β-ionone ring into a keto group, and one of the specific examples is an enzyme activity for synthesizing astaxanthin with zeaxanthin as a substrate by way of 4-ketozeaxanthin (see Fig. 11). In this connection, the polypeptides having such enzyme activities and the DNA strands encoding the polypeptides have not hitherto been reported, and the polypeptides or the DNA strands encoding the polypeptides has no overall homology to polypeptides or DNA strands which have hitherto been reported. Moreover, no such information has been reported that one enzyme has an activity to convert directly a dihydrocarbonyl group of not only the β-ionone ring and the 3-hydroxy-β-ionone ring but also the other compounds into a keto group. Moreover, a homology of CrtW as high as 83% identity at an amino acid sequence level was shown between Agrobacterium and Alcaligenes.

On the other hand, it is possible to allow a microorganisms such as <u>Escherichia coli</u> or the like to produce β-carotene or zeaxanthin by using the carotenoid synthesis genes of the non-photosynthetic bacterium <u>Erwinia</u>, that is the <u>crtE</u>, <u>crtB</u>, <u>crtI</u> and <u>crtY</u> genes of <u>Erwinia</u> afford the microorganism such as <u>Escherichia coli</u> or the like the β-carotene-producing ability, and the <u>crtE</u>, <u>crtB</u>, <u>crtI</u>, <u>crtY</u> and <u>crtZ</u> genes of <u>Erwinia</u> afford the microorganisms such as <u>Escherichia coli</u> or the like the zeaxanthin-producing ability (see Fig. 10 and Laid-Open Publication of WO91/13078). Thus, the substrate of CrtW is supplied by the <u>crt</u> gene cluster of <u>Erwinia</u>, so that when additional <u>crtW</u> gene is introduced into the microorganism such as <u>Escherichia coli</u> or the like which contains the aforementioned <u>crt</u> gene cluster of <u>Erwinia</u>, the β-carotene-producing microorganism will produce canthaxanthin by way of echinenone, and the zeaxanthin-producing microorganism will produce astaxanthin by way of 4-ketozeaxanthin.

Hydroxyl group-introducing enzyme gene (crtZ)

The DNA strands (19) - (27) are genes encoding a hydroxyl group-introducing enzyme (referred to hereinafter as crtZ). Typical examples of the genes are crtZ genes cloned from the marine bacteria Agrobacterium aurantiacus sp. nov. MK1 or Alcaligenes sp. PC-1, which are the DNA strands comprising the nucleotide sequences encoding the polypeptides having the amino acid sequences C - D in Fig. 2 (amino acid Nos. 1 - 162 in SEQ ID NOS: 2) or C - D in Figs. 15 (amino acid Nos. 1 - 162 in SEQ ID NOS: 6). The crtZ gene product (also referred to hereinafter as CrtZ) has an enzyme activity for adding a hydroxyl group to the 3-carbon atom of the β-ionone ring, and one of the specific examples is an enzyme activity for synthesizing zeaxanthin with use of β-carotene as a substrate by way of β-cryptoxanthin (see Fig. 11). In addition, the crtZ gene product also has an enzyme activity for adding a hydroxyl group to the 3-carbon atom of the 4-keto-β-ionone ring, and one of the specific examples is an enzyme activity for synthesizing astaxanthin

with canthaxanthin as a substrate by way of phoenicoxanthin (see Fig. 11). In this connection, the polypeptide having the latter enzyme activity and the DNA strand encoding the polypeptide have not hitherto been reported. Moreover, CrtZ of Agrobacterium and Alcaligenes showed a high homology with CrtZ of Erwinia uredovora (57% and 58% identity), respectively, at an amino acid sequence level. Also, a high homology of 90% identity at an amino acid sequence level was shown between the CrtZ of Agrobacterium and Alcaligenes.

It has been described above that it is possible to allow a microorganism such as Escherichia coli or the like to produce β-carotene by using the carotenoid synthetic genes of the non-photosynthetic bacterium Erwinia. Moreover, it has been described above that it is possible to allow a microorganism such as Escherichia coli or the like to produce canthaxanthin by adding crtW thereto. Thus, the substrate of CrtZ of Agrobacterium or Alcaligenes is supplied by the crtE, crtB, crtl and crtY genes of Enwinia (production of β-carotene), and the crtW gene of Agrobacterium or Alcaligenes added thereto, so that when the crtZ gene of Agrobacterium or Alcaligenes is introduced into a microorganism such as Escherichia coli or the like containing the crt gene group, the β-carotene-producing microorganism will produce zeaxanthin by way of β-cryptoxanthin, and the canthaxanthin-producing microorganism will produce astaxanthin by way of phoenicoxanthin.

Lycopene-cyclizing enzyme gene (crtY)

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The DNA strand encoding the amino acid sequence substantially from E to F of Figs. 3 and 4 (amino acid Nos. 1-386 in SEQ ID NO: 3) is a gene encoding a lycopene-cyclizing enzyme (referred to hereinafter as crtY). A typical example of the gene is the crtY gene cloned from the marine bacterium Agrobacterium aurantiacus sp. nov. MK1, which is the DNA strand comprising the nucleotide sequence encoding the polypeptide having the amino acid sequence E - F in Figs. 3 and 4. The \underline{crtY} gene product (also referred to hereinafter as \underline{CrtY}) has an enzyme activity for synthesizing β carotene with lycopene as a substrate (see Fig. 11). It is possible to allow a microorganism such as Escherichia coli or the like to produce lycopene by using a carotenoid biosynthesis genes of a non-photosynthetic bacterium Erwinia, that is the crtE, crtB and crtI genes of Erwinia give a microorganism such as Escherichia coli or the like a lycopene biosynthesis ability (see Fig. 10, and Laid-Open Publication of WO91/13078). Thus, the substrate of the CrtY of Agrobacterium is supplied by the crt gene group of Envinia, so that when the crtY of Agrobacterium is introduced into a microorganism such as Escherichia coli or the like containing the crt gene group, it is possible to allow the microorganism to produce β -carotene.

In this connection, the CrtY of Agrobacterium has a significant homology of 44.3% identity to the CrtY of Erwinia uredovora at the amino acid sequence level, and these CrtY enzymes also have the same enzymatic function (see Figs. 10 and 11).

Bacteriological properties of marine bacteria

The marine bacteria Agrobacterium aurantiacus sp. nov. MK1 and Alcaligenes sp. PC-1 as the sources of the xanthophyll synthetic genes show the following bacteriological properties.

(Agrobacterium aurantiacus sp. nov. MK1)

(1) Morphology

Form and size of bacterium: rod, 0.9 $\mu m \times 1.2~\mu m;$

Motility: yes;

Flagellum: peripheric flagellum;

Polymorphism of cell: none;

Sporogenesis: none;

Gram staining: negative.

(2) Growths in culture media

Broth agar plate culture: non-diffusive circular orange colonies having a gloss are formed.

Broth agar slant culture: a non-diffusive orange band having a gloss is formed.

Broth liquid culture: homogeneous growth all over the culture medium with a color in orange.

Broth gelatin stab culture: growth over the surface around the stab pore.

(3) Physiological properties

Reduction of nitrate: positive:

Denitrification reaction: negative;

Formation of indole: negative;

Utilization of citric acid: negative;

Formation of pigments: fat-soluble reddish orange pigment;

Urease activity: negative; Oxidase activity: positive; Catalase activity: positive;

β-Glucosidase activity (esculin degradability): positive;

 β -Galactosidase activity: positive;

Growth range: pH, 5 - 9; temperature, 10 - 40°C:

Behavior towards oxygen: aerobic; Durability to seawater: positive;

O - F test: oxidation;

Anabolic ability of saccharides:

Positive D-glucose, D-mannose, D-galactose, D-fructose, lactose, maltose, sucrose, glycogen, N-acetyl-D-glucosamine,

Negative: L-arabinose, D-mannitol, inositol, L-rhamnose, D-sorbitol;

15 Anabolic ability of organic acids:

Positive lactate;

Negative: citrate, malate, gluconate, caprinate, succinate, adipate;

Anabolic ability of the other organic materials:

Positive inosine, uridine, glucose-1-phosphate, glucose-6-phosphate;

Negative: gelatin, L-arginine, DNA, casein.

(Alcaligenes sp. PC-1)

(1) Morphology

25 Form and size of bacterium: short rod, 1.4 µm;

Motility: yes;

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Flagellum: peripheric flagellum; Polymorphism of cell: none;

Sporogenesis: none; Gram staining: negative.

(2) Growths in culture media

Broth agar plate culture: non-diffusive circular orange colonies having a gloss are formed.

Broth agar slant culture: a non-diffusive orange band having a gloss is formed.

Broth liquid culture: homogeneous growth all over the culture medium with a color in orange.

Broth gelatin stab culture: growth over the surface around the stab pore.

(3) Physiological properties

Formation of pigments: fat-soluble reddish orange pigment;

Oxidase activity: positive;Catalase activity: positive;

Growth range: pH, 5 - 9; temperature, 10 - 40°C:

Behavior towards oxygen: aerobic; Durability to seawater: positive;

45 O - F test: oxidation;

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Degradability of gelatin: negative.

Xanthophyll synthetic gene cluster of the other marine bacteria

It has hitherto been reported that 16 marine bacteria have an ability to synthesize ketocarotenoids such as astaxanthin and the like (Yokoyama, A., Izumida, H., Miki, W., "Marine bacteria produced astaxanthin", 10th International Symposium on Carotenoids, Abstract, CL11-3, 1993). If either of the crt genes of the aforementioned marine bacteria Agrobacterium aurantiacus sp. nov. MK-1 or Alcaligenes sp. PC-1 is used as a probe, the gene cluster playing a role of the biosynthesis of ketocarotenoids such as astaxanthin and the like should be obtained from the other astaxanthin producing marine bacteria by using the homology of the genes. In fact, the present inventors have successfully obtained the crtW and crtZ genes as the strongly hybridizing DNA fragments from the chromosomal DNA of Alcaligenes PC-1 with use of a DNA fragment containing crtW and crtZ of Ag. aurantiacus sp. nov. MK1 as a probe (see Examples as for the details). Furthermore, when Alteromonas SD-402 was selected from the remaining 14 marine bacteria having an astaxanthin synthetic ability and a chromosomal DNA was prepared therewith and subjected to the Southern hybridi-

zation experiment with a DNA fragment containing <u>crtW</u> and <u>crtZ</u> of <u>Ag. aurantiacus</u> sp. nov. MK1, the probe hybridized with the bands derived from the chromosomal DNA of the marine bacteria. The DNA strands according to the present invention also includes a DNA strand which hybridizes with the DNA strands (2), (4), (6), (8), (11), (13), (15), (17), (20), (22), (24) and (26).

Acquisition of DNA strands

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Although one of the methods for obtaining the DNA strand having a nucleotide sequence which encodes the amino acid sequence of each enzyme described above is to chemically synthesize at least a part of the strand length according to the method for synthesizing a nucleic acid, it is believed more preferable than the chemical synthetic method to obtain the DNA strand by using the total DNA having been digested with an appropriate restriction enzyme to prepare a library in Escherichia coli, from which library the DNA strand is obtained by the methods conventionally used in the art of genetic engineering such as a hybridization method with an appropriate probe (see the xanthophyll synthetic gene cluster of the other marine bacteria).

Transformation of an microorganism such as Escherichia coli and gene expression

A variety of xanthophylls can be prepared by introducing the present DNA strands described above into appropriate microorganisms such as bacteria, for example <u>Escherichia coli</u>, <u>Zymomonas mobilis</u> and <u>Agrobacterium tumefaciens</u>, and yeasts, for example <u>Saccharomyces cerivisiae</u>.

The outline for introducing an foreign gene into a preferred microorganism is described below.

The procedure or method for introducing and expressing the foreign gene in a microorganism such as <u>Escherichia</u> coli or the like comprises the ones usually used in the art of genetic engineering in addition to those described below in the present invention and may be carried out according to the procedure or method (see, e.g., "Vectors for Cloning Genes", Methods in Enzymology, 216, p. 469-631, 1992, Academic Press, and "Other Bacterial Systems", Methods in Enzymology, 204, p. 305-636, 1991, Academic Press).

(Escherichia coli)

The method for introducing foreign genes into Escherichia coli includes several efficient methods such as the Hanahan's method and the rubidium method, and the foreign genes may be introduced according to these methods (see, for example, Sambrook, J., Fritsch, E.F., Maniatis, T., "Molecular Cloning - A Laboratory Manual", Cold Spring Harbor Laboratory Press, 1989). While foreign genes in Escherichia coli may be expressed according to the conventional methods (see, for example, "Molecular Cloning - A Laboratory Manual"), the expression can be carried out for example with a vector for Escherichia coli having a lac promoter in the pUC or pBluescript series. The present inventors have used a vector pBluescrip II SK or KS for Escherichia coli having a lac promoter and the like to insert the crtW, crtZ and crtY genes of Agrobacterium aurantiacus sp. nov. MK1 and the crtW and crtZ genes of Alcaligenes sp. PC-1 and allowed to express these genes in Escherichia coli.

o (Yeast)

The method for introducing foreign genes into yeast <u>Saccharomyces cerivisiae</u> includes the methods which have already been established such as the lithium method and the like, and the introduction may be carried out according to these methods (see, for example, Ed. Yuichi Akiyama, compiled by Bio-industry Association, "New Biotechnology of Yeast", published by IGAKU SHUPPAN CENTER). Foreign genes can be expressed in yeast by using a promoter and a terminator such as PGK and GPD to construct an expression cassette in which the foreign gene is inserted between the promoter and the terminator so that transcription is led through, and inserting the expression cassette into a vector such as the YRp system which is a multi-copy vector for yeast having the ARS sequence of the yeast chromosome as the replication origin, the YEp system which is a multi-copy vector for yeast having the replication origin of the 2 μm DNA of yeast, and the YIp system which is a vector for integrating a yeast chromosome having no replication origin of yeast (see "New Biotechnology of Yeast", published by IGAKU SHUPPAN CENTER, ibid.; NIPPON NOGEI-KAGAKU KAI ABC Series "Genetic Engineering for Producing Materials", published by ASAKURA SHOTEN; and Yamano, S., Ishii, T., Nakagawa, M., Ikenaga, H., Misawa, N., "Metabolic Engineering for Production of β-carotene and Iycopene in Saccharomyces cerevisiae", Biosci. Biotech. Biochem., 58, p. 1112-1114, 1994).

Zymomonas mobilis)

Foreign genes can be introduced into an ethanol-producing bacterium <u>Zymomonas mobilis</u> by the conjugal transfer method which is common to Gram-negative bacteria, and the foreign genes can be expressed by using a vector pZA22

for <u>Zymomonas mobilis</u> (see Katsumi Nakamura, "Molecular Breeding of <u>Zymomonas mobilis</u>", Nippon Nogei-Kagaku Kaishi, 63, p. 1016-1018, 1989; and Misawa, N., Yamano, S., Ikanaga, H., "Production of β-Carotene in <u>Zymomonas mobilis</u> and <u>Agrobacterium tumefaciens</u> by Introduction of the Biosynthesis Genes from <u>Erwinia uredovora</u>", Appl. Environ. Microbiol., 57, p.1847-1849, 1991).

(Agrobacterium tumefaciens)

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Foreign genes can be introduced into a plant pathogenic bacterium <u>Agrobacterium tumefaciens</u> by the conjugal transfer method which is common to Gram-negative bacteria, and the foreign genes can be expressed by using a vector pBI121 for a bacterium such as <u>Agrobacterium tumefaciens</u> (see Misawa, N., Yamano, S., Ikenaga, H., "Production of β-Carotene in <u>Zymomonas mobilis</u> and <u>Agrobacterium tumefaciens</u> by Introduction of the Biosynthesis Genes from <u>Erwinia uredovora</u>", Appl. Environ. Microbiol., 57, p. 1847-1849, 1991).

Production of xanthophylls by microorganisms

The gene cluster for the synthesis of ketocarotenoids such as astaxanthin derived from a marine bacterium can be introduced and expressed by the procedure or method described above for introducing and expressing an foreign gene in a microorganism.

Farnesyl pyrophosphate (FPP) is a substrate which is common not only to carotenoids but also to other terpenoids such as sesquiterpenes, triterpenes, sterols, hopanols and the like. In general, microorganisms synthesize terpenoids even if they cannot synthesize carotenoids, so that all of the microorganisms should basically have FPP as an intermediate metabolite. Furthermore, the carotenoid synthesis gene cluster of a non-photosynthetic bacterium Erwinia has an ability to synthesize the substrates of the <u>crt</u> gene products of <u>Agrobacterium aurantiacus</u> sp. nov. MK1 or <u>Alcaligenes sp. PC-1</u> by using FPP as a substrate (see Fig. 10). The present inventors have already confirmed that when the group of crt genes of <u>Erwinia</u> is introduced into not only <u>Escherichia coli</u> but also the aforementioned microorganisms, that is the yeast <u>Saccharomyces cerevisiae</u>, the ethanol producing bacterium <u>Zymomonas mobilis</u>, or the plant pathogenic bacterium <u>Agrobacterium tumefaciens</u>, carotenoids such as β-carotene and the like can be produced, as was expected, by these microorganisms (Yamano, S., Ishii, T., Nakagawa, M., Ikenaga, H., Misawa, N., "Metabolic Engineering for Production of β-Carotene and Lycopene in <u>Saccharomyces cerevisiae</u>", Biosci. Biotech. Biochem., 58, p. 1112-1114, 1994; Misawa, N., Yamano, S., Ikenaga, H., "Production of β-Carotene in Zymomonas mobilis and <u>Agrobacterium tumefaciens</u> by Introduction of the Biosynthetic Genes from <u>Erwinia uredovora</u>", Appl. Environ. Microbiol., 57, p. 1847-1849, 1991; and Japanese Patent Application No. 58786/1991 (Japanese Patent Application No. 53255/1990) by the present inventors: "DNA Strands useful for the Synthesis of Carotenoids").

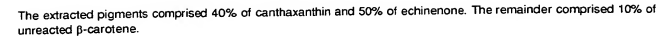
Thus, it should be possible in principle to allow all of the microorganisms, in which the gene introduction and expression system has been established, to produce ketocarotenoids such as astaxanthin and the like by introducing the combination of the carotenoid synthesis gene cluster derived from Enwinia and the DNA strands according to the present invention (typically the carotenoid synthesis gene cluster derived from Agrobacterium aurantiacus sp. nov. MK1 or Alcaligenes sp. PC-1) at the same time into the same microorganism. The process for producing a variety of ketocarotenoids in microorganisms are described below.

(Production of canthaxanthin and echinenone)

It is possible to produce canthaxanthin as a final product and echinenone as an intermediate metabolite by introducing into a microorganism such as Escherichia coli and expressing the crtE, crtB, crtl and crtY genes of Erwinia uredovora required for the synthesis of β-carotene and any one of the DNA strands of the present invention (1) - (9) which is a keto group-introducing enzyme gene (typically, the crtW gene of Agrobacterium aurantiacus sp. nov. MK1 or Alcaligenes PC-1). The yields or the ratio of canthaxanthin and echinenone can be changed by controlling the expression level of the DNA strand (crtW gene) or examining the culturing conditions of a microorganism having the DNA strand. Two embodiments in Escherichia coli are described below, and more details will be illustrated in Examples.

A plasmid pACCAR16AcrtX that a fragment containing the <u>crtE</u>, <u>crtB</u>, <u>crtI</u> and <u>crtY</u> genes of <u>Erwinia uredovora</u> has been inserted into the <u>Escherichia coli</u> vector pACYC184 and a plasmid pAK916 that a fragment containing the <u>crtW</u> gene of <u>Agrobacterium aurantiacus</u> sp. nov. MK1 has been inserted into the <u>Escherichia coli</u> vector pBluescript II SK-were introduced into <u>Escherichia coli</u> JM101 and cultured to the stationary phase to collect bacterial cells and to extract carotenoid pigments. The extracted pigments comprised 94% of canthaxanthin and 6% of echinenone. Also, canthaxanthin was obtained in a yield of 3 mg starting from 2 liters of the culture solution.

A plasmid pACCAR16AcrtX that a fragment containing the <u>crtE</u>, <u>crtB</u>, <u>crtI</u> and <u>crtY</u> genes of <u>Erwinia uredovora</u> has been inserted into the <u>Escherichia coli</u> vector pACYC184 and a plasmid pPC17-3 that a fragment containing the <u>crtW</u> gene of <u>Alcaligenes</u> PC-1 has been inserted into the <u>Escherichia coli</u> vector pBluescript II SK+ were introduced into <u>Escherichia coli</u> JM101 and cultured to the stationary phase to collect bacterial cells and to extract carotenoid pigments.



Production of astaxanthin and 4-ketozeaxanthin>

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It is possible to produce astaxanthin as a final product and 4-ketozeaxanthin as an intermediate metabolite by introducing into a microorganism such as <u>Escherichia coli</u> or the like and expressing the <u>crtE</u>, <u>crtB</u>, <u>crtI</u>, <u>crtY</u> and <u>crtZ</u> genes of <u>Erwinia uredovora</u> required for the synthesis of zeaxanthin and any one of the DNA strands of the present invention . (10) - (18) which is a keto group-introducing enzyme gene (typically, the <u>crtW</u> gene of <u>Agrobacterium aurantiacus</u> sp. nov. MK1 or <u>Alcaligenes</u> PC-1). The yields or the ratio of astaxanthin and 4-ketozeoxanthin can be changed by controlling the expression level of the DNA strand (<u>crtW</u> gene) or examining the culturing conditions of a microorganism having the DNA strand.

Two embodiments in Escherichia coli are described below, and more details will be illustrated in Examples.

A plasmid pACCAR25∆crtX that a fragment containing the <u>crtE</u>, <u>crtB</u>, <u>crtI</u>, <u>crtY</u> and <u>crtZ</u> genes of <u>Erwinia uredovora</u> has been inserted into the <u>Escherichia coli</u> vector pACYC184 and a plasmid pAK916 that a fragment containing the <u>crtW</u> gene of <u>Ag. aurantiacus</u> sp. nov. MK1 has been inserted into the <u>Escherichia coli</u> vector pBluescript II SK- were introduced into <u>Escherichia coli</u> JM101 and cultured to the stationary phase to collect bacterial cells and to extract carotenoid pigments. The yield of the extracted pigments was 1.7 mg of astaxanthin and 1.5 mg of 4-ketozeaxanthin based on 2 liters of the culture solution.

A plasmid pACCAR25AcrtX that a fragment containing the crtE, c

(Production of astaxanthin and phoenicoxanthin)

It is possible to produce astaxanthin as a final product and phoenicoxanthin as an intermediate metabolite by introducing into a microorganism such as <u>Escherichia coli</u> or the like and expressing the <u>crtE</u>, <u>crtB</u>, <u>crtI</u> and <u>crtY</u> genes of <u>Erwinia uredovora</u> required for the synthesis of β-carotene, any one of the DNA strands of the present invention (1) - (9) which is a keto group-introducing enzyme gene (typically, the <u>crtW</u> gene of <u>Agrobacterium aurantiacus</u> sp. nov. MK1 or <u>Alcaligenes</u> PC-1), and any one of the DNA strands of the present invention (19) - (27) which is a hydroxyl group-introducing enzyme gene (typically, the <u>crtZ</u> gene of <u>Ag. aurantiacus</u> sp. nov. MK1 or <u>Alcaligenes</u> PC-1). The yields or the ratio of astaxanthin and phoenicoxanthin can be changed by controlling the expression level of the DNA strands (<u>crtW</u> and <u>crtZ</u> genes) or examining the culturing conditions of a microorganism having the DNA strands. An embodiment in <u>Escherichia coli</u> are described below, and more details will be illustrated in Examples.

A plasmid pACCAR16∆crtX that a fragment containing the <u>crtE</u>, <u>crtB</u>, <u>crtI</u> and <u>crtY</u> genes of <u>Erwinia uredovora</u> has been inserted into the <u>Escherichia coli</u> vector pACYC184 and a plasmid pAK96K that a fragment containing the <u>crtW</u> and <u>crtZ</u> genes of <u>Ag. aurantiacus</u> sp. nov. MK1 has been inserted into the <u>Escherichia coli</u> vector pBluescript II SK-were introduced into <u>Escherichia coli</u> JM101 and cultured to the stationary phase to collect bacterial cells and to extract carotenoid pigments. The yield of the extracted pigments comprised was 3 mg of astaxanthin and 2 mg of phoenicoxanthin starting from 4 liters of the culture solution.

Deposition of microorganisms

Microorganisms as the gene sources of the DNA strands of the present invention and <u>Escherichia coli</u> carrying the isolated genes (the DNA strands of the present invention) have been deposited to National Institute of Bioscience and Human Technology, Agency of Industrial Science and Technology.

(i) Agrobacterium aurantiacus sp. nov. MK1

Deposition No: FERM BP-4506 Entrusted Date: December 20, 1993

(ii) Escherichia coli JM101 (pAccrt-EIB, pAK92)

Deposition No: FERM BP-4505 Entrusted Date: December 20, 1993

(iii) <u>Alcaligense</u> sp. PC-1 Deposition No: FERM BP-4760 Entrusted Date: July 27, 1994

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(iv) Escherichia coli β: pPC17 Deposition No: FERM BP-4761 Entrusted Date: July 27, 1994

Examples

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The present invention is further described more specifically with reference to the following examples without restriction of the invention. In addition, the ordinary experiments of gene manipulation employed herein is based on the standard methods (Sambrook, J., Fritsch, E.F., Maniatis, T., "Molecular Cloning - A Laboratory Manual", Cold Spring Harbor Laboratory Press, 1989), unless otherwise specified.

Example 1: Preparation of chromosomal DNA

Chromosomal DNAs were prepared from three marine bacterial strains, i.e. <u>Agrobacterium aurantiacus</u> sp. nov. MK1, <u>Alcaligenes</u> sp. PC-1, and <u>Alteromonas</u> SD-402 (Yokoyama, A., Izumida, H., Miki, W., "Marine bacteria produced astaxanthin", 10th International Symposium on Carotenoids, Abstract, CL11-3, 1993). After each of these marine bacteria was grown in 200 ml of a culture medium (a culture medium prepared according to the instruction of "Marine Broth" manufactured by DIFCO) at 25°C for 4 days to the stationary phase, the bacterial cells were collected, washed with a TES buffer (20 mM Tris, 10 mM EDTA, 0.1 M NaCl, pH 8), subjected to heat treatment at 68°C for 15 minutes, and suspended into the solution I (50 mM glucose, 25 mM Tris, 10 mM EDTA, pH 8) containing 5 mg/ml of lysozyme (manufactured by SEIKAGAKU KOGYO) and 100 µg/ml of RNase A (manufactured by Sigma). After incubation of the suspension at 37°C for 1 hour, Proteinase K (manufactured by Boehringer-Mannheim) was added and the mixture was incubated at 37°C for 10 minutes. After SARCOSIL (N-lauroylsarcosine Na, manufactured by Sigma) was then added at the final concentration of 1% and the mixture was sufficiently mixed, it was incubated at 37°C for several hours. The mixture was extracted several times with phenol/chloroform, and ethanol in a two-time amount was added slowly. Chromosomal DNA thus deposited was wound around a glass rod, rinsed with 70% ethanol and dissolved in 2 ml of a TE buffer (10 mM Tris, 1 mM EDTA, pH 8) to prepare a chromosomal DNA solution.

Example 2: Preparation of hosts for a cosmid library

(1) Preparation of phytoene-producing Escherichia coli

After the removal of the BstEII (1235) - Eco521 (4926) fragment from a plasmid pCAR16 having a carotenoid synthesis gene cluster except the crtZ gene of Erwinia uredovora (Misawa, N.,Nakagawa, M., Kobayashi, K., Yamano, S., Izawa, Y., Nakamura, K., Harashima, K., "Elucidation of the Erwinia uredovora Carotenoid Biosynthetic Pathway by Functional Analysis of Gene Porducts expressed in Escherichia coli", J. Bacteriol., 172, p. 6704-6712, 1990; and Japanese Patent Application No. 58786/1991 (Japanese Patent Application No. 53255/1990): "DNA Strands useful for the Synthesis of Carotenoids"), a 2.3 kb Asp718 (KpnI) - EcoRI fragment containing the crtE and crtB genes required for the production of phytoenes was cut out. This fragment was then inserted into the EcoRV site of the E. coli vector pACYC184 to give an aimed plasmid (pACCRT-EB). The bacterium E. coli containing pACCRT-EB exhibits resistance to an antibiotic chloramphenicol (Cm¹) and produces phytoenes (Linden, H., Misawa, N., Chamovitz, D., Pecker, I., Hirschberg, J., Sandmann, G., "Functional Complementation in Escherichia coli of Different Phytoene Desaturase Genes and Analysis of Accumulated Carotenes", Z. Naturforsch., 46c, 1045-1051, 1991).

5 (2) Preparation of lycopene-producing Escherichia coli

After the removal of the BstEII (1235) - SnaBI (3497) fragment from a plasmid pCAR16 having a carotenoid synthesis gene cluster except the crtZ gene of Erwinia uredovora, a 3.75 kb Asp718 (KpnI) - EcoRI fragment containing the crtE, crtI and crtB genes required for the production of lycopene was cut out. This fragment was then inserted into the EcoRV site of the E. coli vector pACYC184 to give an aimed plasmid (pACCRT-EIB). The bacterium E. coli containing pACCRT-EIB exhibits Cmr and produces lycopene (Cunningham Jr, F.X., Chamovitz, D., Misawa, N., Gatt, E., Hirschberg, J., "Cloning and Functional Expression in Escherichia coli of Cyanobacterial Gene for Lycopene Cyclase, the Enzyme that catalyzes the Biosynthesis of β-Carotenes", FEBS Lett., 328, 130-138, 1993).

(3) Preparation of β-carotene-producing Escherichia coli

After the <u>crtX</u> gene was inactivated by subjecting a plasmid pCAR16 having a carotenoid synthesis gene cluster except the <u>crtZ</u> gene of <u>Erwinia uredovora</u> to digestion with restriction enzyme <u>BstEII</u>, the Klenow fragment treatment and the ligation reaction, a 6.0 kb <u>Asp</u>718 (<u>KpnI</u>) - <u>EcoRI</u> fragment containing <u>crtE</u>, <u>crtY</u>, <u>crtI</u> and <u>crtB</u> genes required

for the production of β -carotene was cut out. This fragment was then inserted into the <u>EcoRV</u> site of the <u>E. coli</u> vector pACYC184 to give an aimed plasmid (referred to hereinafter as pACCAR16 Δ crtX). The bacterium <u>E. coli</u> containing pACCAR16 Δ crtX exhibits Cm^r and produces β -carotene. In this connection, the restriction enzyme and enzymes used for genetic manipulation have been purchased from TAKARA SHUZO (K.K.) or Boehringer-Mannheim.

Example 3: Preparation of a cosmid library and acquisition of Escherichia coli which exhibits orange in color

After the restriction enzyme <u>Sau</u>3Al was added in an amount of one unit to 25 µg of the chromosomal DNA of <u>Agrobacterium aurantiacus</u> sp. nov. MK1, the mixture was incubated at 37°C for 15 minutes and heat treated at 68°C for 10 minutes to inactivate the restriction enzyme. Under the condition, many partially digested fragments with <u>Sau</u>3Al were obtained at about 40 kb. The cosmid vector pJBB (resistant to ampicillin (Ap^r)) which had been subjected to <u>Bam</u>HI digestion and alkaline phosphatase treatment and the right arm (shorter fragment) of pJBB which had been digested with <u>Sall/Bam</u>HI and then recovered from the gel were mixed with a part of the above <u>Sau</u>3Al partial fragments, and ligated at 12°C overnight. In this connection, pJBB has been purchased from Amersham.

Phage particles were obtained in an amount sufficient for preparing a cosmid library by the <u>in vitro</u> packaging with a Gigapack Gold (manufactured by Stratagene; available from Funakoshi) using the DNA above ligated.

After Escherichia coli DH1 (ATCC33849) and Escherichia coli DH1, each of which has one of the three plasmids prepared in Example 2, were infected with the phage particles, these bacteria were diluted so that 100 - 300 colonies were found on a plate, plated on LB containing appropriate antibiotics (1% trypton, 0.5% yeast extract, 1% NaCl), and cultured at 37°C or room temperature for a period of overnight to several days.

As a result, in cosmid libraries having the simple Escherichia coli (beige) or the phytoene-producing Escherichia coli (beige) with pACCRT-EB as a host, no colonies with changed color were obtained notwithstanding the screening of a ten thousand or more of the colonies for respective libraries. On the other hand, in cosmid libraries having the lycopene-producing Escherichia coli (light red) with pACCRT-EIB or the β-carotene-producing Escherichia coli (yellow) with pACCAR16ΔcrtX as a host, colonies exhibiting orange have appeared in a proportion of one strain to several hundred colonies, respectively. Most of these transformed Escherichia coli strains which exhibits orange contained plasmid pJB8 in which about 40 kb partially digested Sau3AI fragments were cloned. It is also understood from the fact that no colonies with changed color appeared in cosmid libraries having the simple Escherichia coli or the phytoene-producing Escherichia coli with pACCRT-EB as a host, that Escherichia coli having an ability of producing a carotenoid synthetic intermediate of the later steps of at least phytoene should be used as a host for the purpose of expression-cloning the xanthophyll synthesis gene cluster from the chromosomal DNA of Agrobacterium aurantiacus sp. nov. MK1.

Example 4: Localization of a fragment containing an orange pigment synthesis gene cluster

When individual several ten colonies out of the orange colonies obtained in cosmid libraries having the lycopene-producing Escherichia coli (light red) with pACCRT-EIB or the β-carotene-producing Escherichia coli (yellow) with pACCRT6ΔcrtX as a host were selected to analyze the plasmids, 33 kb - 47 kb fragments partially digested with Sau3Al were inserted in vector pJB8 in all of the colonies except one strain. The remaining one strain (lycopene-producing Escherichia coli as a host) contains a plasmid, in which a 3.9 kb fragment partially digested with Sau3Al was inserted in pJB8 (referred to hereinafter as plasmid pAK9). This was considered to be the one formed by the in vivo deletion of the inserted fragment after the infection to Escherichia coli. The same pigment (identified as astaxanthin in Example 6) as that in the orange colonies obtained from the other cosmid libraries was successfully synthesized with the lycopene-producing Escherichia coli having pAK9, pAK9 was used as a material in the following analyses.

Example 5: Determination of the nucleotide sequence in the orange pigment synthesis gene cluster

A 3.9 kb EcoRI inserted fragment prepared from pAK9 was inserted into the EcoRI site of the Escherichia coli vector pBluescrip II SK+ to give two plasmids (pAK91 and pAK92) with the opposite directions of the fragment to the vector. The restriction enzyme map of one of the plasmids (pAK92) is illustrated in Fig. 12. When pAK92 was introduced into the lycopene-producing Escherichia coli, orange colonies were obtained as a result of the synthesis of astaxanthin (Example 6). However, no ability for synthesizing new pigments was afforded even if pAK91 was introduced into the lycopene-producing Escherichia coli. It was thus considered that the pigment synthesis gene cluster in the plasmid pAK92 has the same direction as that of the lac promoter of the vector. Next, each of a 2.7 kb Pstl fragment obtained by the Pstl digestion of pAK91, a 2.9 kb BamHI fragment obtained by the BamHI digestion of pAK92, and 2.3 kb and 1.6 kb Sall fragments obtained by the Sall digestion of pAK92 was cloned into the vector pBluescrip II SK-. The restriction maps of plasmids referred to as pAK94, pAK96, pAK98, pAK910, pAK93, and pAK95 are illustrated in Fig. 12. The plasmids pAK94, pAK96, pAK98 and pAK910 have the pigment synthesis gene cluster in the same direction as that of the lac promoter of the vector, while the plasmids pAK93 and pAK95 have the pigment synthesis gene cluster in the opposite direction to that of the promoter.

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It was found that when the plasmid pAK96 having a 2.9 kb <u>Bam</u>HI fragment was introduced into the lycopene-producing <u>Escherichia coli</u>, the transformant also synthesized astaxanthin as in the case when the plasmid pAK92 having a 3.9 kb <u>Eco</u>RI fragment was introduced (Example 6), so that the DNA sequence of the 2.9 kb <u>Bam</u>HI fragment was determined

The DNA sequence was determined by preparing deletion mutants of the 2.9 kb BamHI fragment from the normal and opposite directions and determining the sequence using clones having various lengths of deletions. The deletion mutants were prepared from the four plasmids pAK96, pAK98, pAK93 and pAK95 according to the following procedure: Each of the plasmids, 10 μg, was decomposed with Sacl and Xbal and extracted with phenol/chloroform to recover DNA by ethanol precipitation. Each of DNA was dissolved in 100 µl of ExoIII buffer (50 mM Tris-HCl, 100 mM NaCl, 5 mM MgCl₂, 10 mM 2-mercaptoethanol, pH 8.0), 180 units of ExoIII nuclease was added, and the mixture was maintained at 37°C. A 10 µl portion was sampled at every 1 minute, and two samples were transferred into a tube in which 20 µl of MB buffer (40 mM sodium acetate, 100 mM NaCl, 2 mM ZnCl₂, 10% glycerol, pH 4.5) is contained and which is placed on ice. After completion of the sampling, five tubes thus obtained were maintained at 65°C for 10 minutes to inactivate the enzyme, five units of mung bean nuclease were added, and the mixtures were maintained at 37°C for 30 minutes. After the reaction, five DNA fragments different from each other in the degrees of deletion were recovered for each plasmid by agarose gel electrophoresis. The DNA fragments thus recovered was blunt ended with the Klenow fragment, subjected to the ligation reaction at 16°C overnight, and Escherichia coli JM109 was transformed. A single stranded DNA was prepared from each of various clones thus obtained with a helper phage M13K07, and subjected to the sequence reaction with a fluorescent primer cycle-sequence kit available from Applied Biosystem (K.K.), and the DNA sequence was determined with an automatic sequencer.

The DNA sequence comprising 2886 base pairs (bp) thus obtained is illustrated in Figs. 5 - 9 (SEQ ID NO: 4). As a result of examining an open reading frame having a ribosome binding site in front of the initiation codon, three open reading frames which can encode the corresponding proteins (A - B (nucleotide positions 229 - 864 of SEQ ID NO: 4), C - D (nucleotide positions 864 - 1349), E - F (nucleotide positions 1349 - 2506) in Figs. 5 - 9) were found at the positions where the three xanthophyll synthesis genes <u>crtW</u>, <u>crtZ</u> and <u>crtY</u> are expected to be present. For the two open reading frames of A - B and E - F, the initiating codon is GTG, and for the remaining open reading frame C - D, it is ATG.

Example 6: Identification of the orange pigment

The lycopene-producing Escherichia coli JM101 having pAK92 or pAK96 introduced thereinto (Escherichia coli (pACCRT-EIB, pAK92 or pAK96); exhibiting orange) or the β-carotene-producing Escherichia coli JM101 having pAK94 or pAK96K (Fig. 12) introduced thereinto (Escherichia coli (pACCAR16ΔcrtX, pAK94 or pAK96K); exhibiting orange) was cultured in 4 liters of a 2YT culture medium (1.6% trypton, 1% yeast extract, 0.5% NaCl) containing 150 μg/ml of ampicillin (Ap, manufactured by Meiji Seika) and 30 μg/ml of chloramphenicol (Cm, manufactured by Sankyo) at 37°C for 18 hours. Bacterial cells collected from the culture solution was extracted with 600 ml of acetone, concentrated, extracted twice with 400 ml of chloroform/methanol (9/1), and concentrated to dryness. Then, thin layer chromatography (TLC) was conducted by dissolving the residue in a small amount of chloroform/methanol (9/1) and developing on a silica gel plate for preparative TLC manufactured by Merck with chloroform/methanol (15/1). The original orange pigment was separated into three spots at the Rf values of 0.72, 0.82 and 0.91 by TLC. The pigment of the darkest spot at Rf 0.72 corresponding to 50% of the total amount of orange pigment and the pigment of secondly darker spot at Rf 0.82 were scratched off from the TLC plate, dissolved in a small amount of chloroform/methanol (9/1) or methanol, and chromatographed on a Sephadex LH-20 column (15 × 300 mm) with an eluent of chloroform/methanol (9/1) or methanol to give purified materials in a yield of 3 mg (Rf 0.72) and 2 mg (Rf 0.82), respectively.

It has been elucidated from the results of the UV-visible, ¹H-NMR and FD-MS (m/e 596) spectra that the pigment at Rf 0.72 has the same planar structure as that of astaxanthin. When the pigment was dissolved in diethyl ether: 2-propanol: ethanol (5:5:2) to measure the CD spectrum, it was proved to have stereochemical configuration of 3S, 3'S, and thus identified as astaxanthin; see Fig. 11 for the structural formula). Also, the pigment at Rf 0.82 was identified as phoenicoxanthin (see Fig. 11 for the structural formula) from the results of its UV-visible, ¹H-NMR and FD-MS (m/e 580) spectra. In addition, the pigment at 0.91 was canthaxanthin (Example 7(2)).

Example 7: Identification of metabolic intermediates of xanthophyll

(1) Identification of 4-ketozeaxanthin

The zeaxanthin producing <u>Escherichia coli</u> was prepared according to the following procedure. That is to say, the plasmid pCAR25 having total carotenoid synthesis gene cluster of <u>Er. uredorora</u> (Misawa, N., Nakagawa, M., Kobayashi, K., Yamano, S., Izawa, Y., Nakamura, K., Harashima, K., "Elucidation of the <u>Erwinia uredovora</u> Carotenoid Biosynthetic Pathway by Functional Analysis of Gene Products expressed in <u>Escherichia coli</u>", J. Bacteriol., 172, p. 6704-6712, 1990; and Japanese Patent Application No. 58786/1991 (Japanese Patent Application No. 53255/1990): "DNA

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Strands useful for the Synthesis of Carotenoids") was digested with restriction enzyme <u>Bst</u>EII, and subjected to the Klenow fragment treatment and ligation reation to inactivate the <u>crtX</u> gene by reading frame shift, and then a 6.5 kb <u>Asp</u>718 (<u>KpnI</u>) - <u>EcoRI</u> fragment containing the <u>crtE</u>, <u>crtY</u>, <u>crtI</u>, <u>crtB</u> and <u>crtZ</u> genes required for producing zeaxanthin was cut out. This fragment was then inserted into the <u>EcoRV</u> site of the <u>Escherichia coli</u> vector pACYC184 to give the aimed plasmid (referred to hereinafter as pACCAR25∆crtX).

The zeaxanthin-producing Escherichia coli JM101 having pAK910 or pAK916 (Fig. 12) introduced thereinto (Escherichia coli (pACCAR25ΔcrtX, pAK910 or pAK916); exhibiting orange) was cultured in 2 liters of a 2YT culture medium containing 150 μg/ml of Ap and 30 μg/ml of Cm at 37°C for 18 hours. Bacterial cells collected from the culture solution was extracted with 300 ml of acetone, concentrated, extracted twice with 200 ml of chloroform/methanol (9/1), and concentrated to dryness. Then, thin layer chromatography (TLC) was conducted by dissolving the residue in a small amount of chloroform/methanol (9/1) and developing on a silica gel plate for preparative TLC manufactured by Merck with chloroform/methanol (15/1). The original orange pigment was separated into three spots at the Rf values of 0.54 (46%). 0.72 (53%) and 0.91 (1%) by TLC. The pigment at Rf 0.54 was scratched off from the TLC plate, dissolved in a small amount of chloroform/methanol (9/1) or methanol, and chromatographed on a Sephadex LH-20 column (15 × 300 mm) with an eluent of chloroform/methanol (9/1) or methanol to give a purified material in a yield of 1.5 mg.

This material was identified as 4-ketozeaxanthin (see Fig. 11 for the structural formula) since its UV-visible spectrum, FD-MS spectrum (m/e 582) and mobility in silica gel TLC (developed with chloroform/methanol (15/1)) accorded perfectly with those of the standard sample of 4-ketozeaxanthin (purified from Agrobacterium aurantiacus sp. nov. MK1; Japanese Patent Application No. 70335/1993). In addition, the pigments at Rf 0.72 and 0.91 are astaxanthin (Example 6) and canthaxanthin (Example 7 (2)), respectively.

(2) Identification of canthaxanthin

The β -carotene producing Escherichia coli JM101 having pAK910 or pAK916 introduced thereinto (Escherichia coli (pACCAR16 Δ crtX, pAK910 or pAK916); exhibiting orange) was cultured in 2 liters of a 2YT culture medium containing 150 μ g/ml of Ap and 30 μ g/ml of Cm at 37°C for 18 hours. Bacterial cells collected from the culture solution was extracted with 300 ml of acetone, concentrated, extracted twice with 200 ml of chloroform/methanol (9/1), and concentrated to dryness. Then, thin layer chromatography (TLC) was conducted by dissolving the residue in a small amount of chloroform/methanol (9/1) and developing on a silica gel plate for preparative TLC manufactured by Merck with chloroform/methanol (50/1). The pigment of the darkest spot corresponding to 94% of the total amount of orange pigments was scratched off from the TLC plate, dissolved in a small amount of chloroform/methanol (9/1) or chloroform/methanol (1/1), and chromatographed on a Sephadex LH-20 column (15 × 300 mm) with an eluent of chloroform/methanol (9/1) or chloroform/methanol (1/1) to give a purified material in a yield of 3 mg.

This material was identified as canthaxanthin (see Fig. 11 for the structural formula) since its UV-visible, 1 H-NMR, FD-MS (m/e 564) spectra and mobility in silica gel TLC (Rf 0.53 on developing with chloroform/methanol (50/1)) accorded perfectly with those of the standard sample of canthaxanthin (manufactured by BASF). In addition, the pigment corresponding to 6% of the total orange pigments found in the initial extract was considered echinenone (see Fig. 11 for the structural formula) on the basis of its UV-visible spectrum, mobility in silica gel TLC (Rf 0.78 on developing with chloroform/methanol (50/1)), and mobility in HPLC with NOVA PACK HR 6 μ C18 (3.9 × 300 mm; manufactured by Waters) (RT 16 minutes on developing at a flow rate of 1.0 ml/min with acetonitrile/methanol/2-propanol (90/6/4)).

(3) Identification of zeaxanthin

The β-carotene-producing Escherichia coli JM101 having pAK96NK introduced thereinto (Escherichia coli (pACCAR16ΔcrtX, pAK96NK); exhibiting yellow) was cultured in 2 liters of a 2YT culture medium containing 150 μg/ml of Ap and 30 μg/ml of Cm at 37°C for 18 hours. Bacterial cells collected from the culture solution was extracted with 300 ml of acetone, concentrated, extracted twice with 200 ml of chloroform/methanol (9/1), and concentrated to dryness. Then, thin layer chromatography (TLC) was conducted by dissolving the residue in a small amount of chloroform/methanol (9/1) and developing on a silica gel plate for preparative TLC manufactured by Merck with chloroform/methanol (9/1). The pigment of the darkest spot corresponding to 87% of the total amount of yellow pigments was scratched off from the TLC plate, dissolved in a small amount of chloroform/methanol (9/1) or methanol, and chromatographed on a Sephadex LH-20 column (15 × 300 mm) with an eluent of chloroform/methanol (9/1) or methanol to give a purified material in a yield of 3 mg.

It has been elucidated that this material has the same planar structure as that of zeaxanthin since its UV-visible, ¹H-NMR, FD-MS (m/e 568) spectra and mobility in silica gel TLC (Rf 0.59 on developing with chloroform/methanol (9/1)) accorded perfectly with those of the standard sample of zeaxanthin (manufactured by BASF). When the pigment was dissolved in diethyl ether: 2-propanol: ethanol (5:5:2) to measure the CD spectrum, it was proved to have a stereochemical configuration of 3R, 3'R, and thus identified as zeaxanthin (see Fig. 11 for the structural formula). Also, the pigment corresponding to 13% of the total yellow pigments found in the initial extract was considered β-cryptoxan-

thin (see Fig. 11 for the structural formula) on the basis of its UV-visible spectrum, mobility in silica gel TLC (Rf 0.80 on developing with chloroform/methanol (9/1)), and mobility in HPLC with NOVA PACK HR 6μ C18 (3.9 × 300 mm; manufactured by Waters) (RT 19 minutes on developing at a flow rate of 1.0 ml/min with acetonitrile/methanol/2-propanol (90/6/4)).

(4) Identification of β -carotene

The lycopene-producing Escherichia coli JM101 having pAK98 introduced thereinto (Escherichia coli (pACCRT-EIB, pAK98); exhibiting yellow) was cultured in 2 liters of a 2YT culture medium containing 150 μ g/ml of Ap and 30 μ g/ml of Cm at 37°C for 18 hours. Bacterial cells collected from the culture solution was extracted with 300 ml of acetone, concentrated, and extracted twice with 200 ml of hexane. The hexane layer was concentrated and chromatographed on a silica gel column (15 \times 300 mm) with an eluent of hexane/ethyl acetate (50/1) to give 3 mg of a purified material.

The material was identified as β -carotene (see Fig. 11 for the structural formula), since all of the data of its UV-visible, FD-MS spectrum (m/e 536) and mobility in HPLC with NOVA PACK HR 6μ C18 (3.9 × 300 mm; manufactured by Waters) (RT 62 minutes on developing at a flow rate of 1.0 ml/min with acetonitrile/methanol/2-propanol (90/6/4)) accorded with those of the standard sample of β -carotene (all trans type; manufactured by Sigma).

Example 8: Identification of xanthophyll synthesis gene cluster

(1) Identification of a keto group-introducing enzyme gene

It is apparent from the results of Example 6 that among the 3.9 kb fragment contained in pAK9 (Example 4) or pAK92, all of the genes required for the synthesis of astaxanthin from lycopene is contained in the 2.9 kb BamHI fragment at the right side (pAK96, Fig. 12). Thus, the 1.0 kb fragment at the left side is not needed. Unique Ncol and Kpnl sites are present within the 2.9 kb BamHI fragment of pAK96. It is found from the results of Example 7 (3) that the 1.4 kb fragment (pAK96NK) between the Ncol and Kpnl sites has a hydroxyl group-introducing enzyme activity but has no keto group-introducing enzyme activity. Canthaxanthin can also be synthesized from β-carotene with the 2.9 kb BamHI fragment from which a fragment of the right side from unique Sall site between the Ncol and Konl sites had been removed (pAK910) or with the 2.9 kb BamHI fragment from which a fragment of the right side from the HinclI site positioned at the left side of the Sall site had been removed (pAK916), but activity for synthesizing canthaxanthin from βcarotene disappeared in the 2.9 kb BamHI fragment of pAK96 from which a fragment of the right side from the Ncol site left of the Hincll site had been removed. On the other hand, even if a fragment of the left side from unique BallI site which is present leftward within the 0.9 kb BamHI - HincII fragment of pAK916 was removed, similar activity to that of the aforementioned BamHI - HincII fragment (pAK916) was observed. It is thus considered that a gene encoding a keto group-introducing enzyme having an enzyme activity for synthesizing canthaxanthin from β -carotene as a substrate is present within the 0.74 kb BgIII - HincII fragment of pAK916, and the aforementioned Ncol site is present within this gene. As a result of determining the nucleotide sequence, an open reading frame which corresponds to the gene and has a ribosome binding site just in front of the initiation codon was successfully detected, it was then designated as the crtW gene. The nucleotide sequence of the crtW gene and the encoded amino acid sequence are illustrated in Fig. 1 (SEQ ID NO: 1).

The $\underline{\operatorname{crtW}}$ gene product (CrtW) of $\underline{\operatorname{Agrobacterium aurantiacus}}$ sp. nov. MK1 has an enzyme activity for converting a methylene group at the 4-position of a β -ionone ring into a keto group, and one of the specific examples is an enzyme activity for synthesizing canthaxanthin from β -carotene as a substrate by way of echinenone (Example 7 (2); see Fig. 11). Furthermore, the $\underline{\operatorname{crtW}}$ gene product also has an enzyme activity for converting a methylene group at the 4-position of a 3-hydroxy- β -ionone ring into a keto group, and one of the specific examples is an enzyme activity for synthesizing astaxanthin from zeaxanthin as a substrate by way of 4-ketozeaxanthin (Example 7 (1); see Fig. 11). In addition, polypeptides having such enzyme activities and DNA strands encoding these polypeptides have not hitherto been known, and the polypeptides and the DNA strands encoding these polypeptides have no overall homology to any polypeptides or DNA strands having been hitherto known. Also, no such informations have hitherto been described that a methylene group of not only a β -ionone ring and a 3-hydroxy- β -ionone ring but also the other compounds is directly converted into a keto group with an enzyme.

(2) Identification of a hydroxyl group-introducing enzyme gene

Unique <u>Sal</u>I site is present within the 2.9 kb <u>Bam</u>HI fragment of pAK96. When the 2.9 kb <u>Bam</u>HI fragment is cut into two fragments at the <u>Sal</u>I site, these two fragments (pAK910 and pAK98) have no hydroxyl group-introducing activity. That is to say, the left fragment (pAK910) has only a keto group-introducing enzyme activity (Example 7 (2)), and the right fragment (pAK98) has only a lycopene-cyclizing enzyme activity (Example 7 (4)). On the other hand, when a 1.4 kb $N\infty$ I - KpnI fragment (pAK96NK) containing the aforementioned <u>Sal</u>I site is introduced into a β -carotene-producing

Escherichia coli, zeaxanthin is synthesized by way of β-cryptoxanthin (Example 7 (3)). It is thus considered that a gene encoding a hydroxyl group-introducing enzyme which has an enzyme activity for synthesizing zeaxanthin from β-carotene as a substrate is present within the 1.4 kb Ncol - Kpnl fragment of pAK96NK, and the aforementioned Sall site is present within this gene. As a result of determining the nucleotide sequence, an open reading frame which corresponds to the gene and has a ribosome binding site just in front of the initiation codon was successfully detected, it was then referred to as the crtZ gene. The nucleotide sequence of the crtZ gene and the encoded amino acid sequence are illustrated in Fig. 2 (SEQ ID NO: 2).

The \underline{crtZ} gene product (CrtZ) of $\underline{Agrobacterium\ aurantiacus}$ sp. nov. MK1 has an enzyme activity for adding a hydroxyl group to the 3-carbon of a β -ionone ring, and one of the specific examples is an enzyme activity for synthesizing zeaxanthin from β -carotene as a substrate by way of β -cryptoxanthin (Example 7 (3); see Fig. 11). Furthermore, the \underline{crtZ} gene product also has an enzyme activity for adding a hydroxyl group to the 3-carbon of a 4-keto- β -ionone ring, and one of the specific examples is an enzyme activity for synthesizing astaxanthin from canthaxanthin as a substrate by way of phoenicoxanthin (Example 6; see Fig. 11). In addition, polypeptides having the latter enzyme activity and DNA strands encoding these polypeptides have not hitherto been known. Also, the CrtZ of $\underline{Agrobacterium}$ showed significant homology to the CrtZ of $\underline{Erwinia\ uredovora}$ (identity of 57%) at the level of amino acid sequence.

(3) Identification of a lycopene cyclase gene

Astaxanthin can be synthesized from β-carotene with the 2.9 kb BamHI fragment from which a fragment of the right side from a KpnI site had been removed (pAK96K) or with the 2.9 kb BamHI fragment from which a fragment right from the PstI site which is placed further right of the KpnI site had been removed (pAK94) (Example 6), but astaxanthin cannot be synthesized from lycopene. On the other hand, when a 1.6 kb Sall fragment (pAK98), which contains a right fragment from unique Sall site present further left than the aforementioned KpnI site within the 2.9 kb BamHI fragment, was introduced into lycopene-producing Escherichia coli. β-carotene was synthesized (Example 7 (4)). It is thus considered that a gene encoding lycopene cyclase that has an enzyme activity for synthesizing β-carotene from lycopene as a substrate is present within the 1.6 kb Sall fragment of pAK98, and this gene is present over a range of the KpnI site and the PstI site. As a result of determining the nucleotide sequence, an open reading frame which corresponds to the gene and has a ribosome binding site just in front of the initiation codon was successfully detected, it was then referred to as the crtY gene. The nucleotide sequence of the crtY gene and the amino acid sequence to be encoded are illustrated in Figs. 3 - 4 (SEQ ID NO: 3).

The <u>crtY</u> gene product (CrtY) of <u>Agrobacterium aurantiacus</u> sp. nov. MK1 has significant homology to the CrtY of <u>Erwinia uredovora</u> (identity of 44.3%) at the level of amino acid sequence, and the functions of both enzymes are the same.

Example 9: Southern blotting analysis with the chromosomal DNA of the other marine bacteria

Examination was conducted whether a region exhibiting homology with the isolated <u>crtW</u> and <u>crtZ</u> is obtained from a chromosomal DNAs of the other marine microorganisms. The chromosomal DNAs of <u>Alcaligenes</u> sp. PC-1 and <u>Alteromonas</u> sp. SD-402 prepared in Example 1 were digested with restriction enzymes <u>BamHI</u> and <u>PstI</u>, and separated by agarose gel electrophoresis. All of the DNA fragments thus separated were denaturated with an alkali solution of 0.5 N NaOH and 1.5 M NaCl, and transferred on a nylon membrane filter over an overnight period. The nylon membrane filter on which DNAs had been adsorbed was dipped in a hybridization solution (6 × Denhardt, 5 × SSC, 100 μg/ml ssDNA), and pre-hybridization was conducted at 60°C for 2 hours. Next, the 1.5 kb DNA fragment cut out from pAK96K with <u>Ball</u>, which contains <u>crtW</u> and <u>crtY</u>, was labelled with a Mega primeTM DNA labelling systems (Amersham) and [α-32P]dCTP (~110TBq/mmol) and added to the aforementioned prehybridization solution to conduct hybridization at 60°C for 16 hours.

After hybridization, the filter was washed with $2 \times SSC$ containing 0.1% SDS at $60^{\circ}C$ for 1 hour, and subjected to the detection of signals showing homology by autoradiography. As a result, strong signals were obtained at about 13 kb in the product digested with $\underline{Bam}HI$ and at 2.35 kb in the product digested with $\underline{Pst}I$ in the case of $\underline{Alcaligenes}$ sp. PC-1, and strong signals were obtained at about 5.6 kb in the product digested with $\underline{Bam}HI$ and at 20 kb or more in the product digested with $\underline{Pst}I$ in the case of $\underline{Alteromonas}$ sp. SD-4.

Example 10: Acquisition of a xanthophyll synthesis gene cluster from the other marine bacterium

As it was found from the results of Example 9 that the <u>Pst</u>I digest of the chromosomal DNA of <u>Alcaligenes</u> sp. PC-1 has a region of about 2.35 kb hybridizing with a DNA fragment containing the <u>crtW</u> and <u>crtZ</u> genes of <u>Agrobacterium aurantiacus</u> sp. nov. MK1, the chromosomal DNA of <u>Alcaligenes</u> was digested with <u>Pst</u>I, and then DNA fragments of 2 - 3.5 kb in size was recovered by agarose gel electrophoresis. The DNA fragments thus collected were inserted into the <u>Pst</u>I site of a vector pBluescript II SK+, and introduced into <u>Escherichia coli</u> DH5α to prepare a partial library of <u>Alcaligenes</u>

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genes. When the partial library was subjected to colony hybridization with a 1.5 kb DNA fragment containing the crtW and crtZ genes of Agrobacterium as a probe, a positive colony was isolated from about 5,000 colonies. In this case, colony hybridization was conducted under the same condition as in the Southern blotting analysis shown in Example 9. When plasmid DNA was isolated from the colony thus obtained, and digested with Pst to examine the size of the integrated DNA fragments, it was found that the plasmid contained three different fragments. Thus, a 2.35 kb fragment to be hybridized was selected from the three different DNA fragments by the Southern blotting analysis described in Example 9, the 2.35 kb Pst fragment was recovered by agarose gel electrophoresis and inserted again into the Pst site of pBluescript II SK+ to prepare the plasmids pPC11 and pPC12. In pPC11 and pPC12, the aforementioned 2.35 kb Pst fragment was inserted into the Pst I site of pBluescript II SK+ in an opposite direction to each other. The restriction enzyme map of pPC11 is illustrated in Fig. 19.

Example 11: Determination of nucleotide sequence of xanthophyll synthesis gene cluster in Alcaligenes

When each of pPC11 and pPC12 was introduced into β-carotene-producing Escherichia coli, orange colonies were obtained due to the synthesis of astaxanthin (Example 12) in the former, but no other pigments were newly synthesized in the latter. It was thus considered that the direction of the astaxanthin synthesis gene cluster in the plasmid pPC11 was the same as that of the vector lac promoter. It was also found that pPC11 contained no lycopene cyclizing enzyme genes, since no other pigments were newly produced even if pPC11 was introduced into the lycopene-producing Escherichia coli.

It was found that even if a plasmid having a 0.72 kb <u>Bst</u>Ell - <u>Eco</u>RV fragment positioned at the right side of the <u>Pst</u>I fragment had been removed (referred to as pPC17, Fig. 19) was introduced into the β-carotene-producing <u>Escherichia coli</u>, the transformant of <u>Escherichia coli</u> synthesized astaxanthin and the like (Example 12), same as in the case of <u>E. coli</u> into which pPC11 was introduced, so that the nucleotide sequence of the 1.63 kb <u>Pst</u>I - <u>Bst</u>Ell fragment in pPC17 was determined.

Deletion mutants were prepared with pPC17 and pPC12 according to the following procedure. A 10 μg portion of each of pPC17 and pPC12 was digested with <u>KpnI</u> and <u>HindIII</u> or <u>KpnI</u> and <u>Eco</u>RI, extracted with phenol/chloroform, and DNA was recovered by precipitation with ethanol. Each of DNAs was dissolved in 100 μl of <u>Exo</u>III buffer (50 mM Tris-HCI, 100 mM NaCI, 5 mM MgCl₂, 10 mM 2-mercaptoethanol, pH 8.0), 180 units of <u>Exo</u>III nuclease was added, and the mixture was maintained at 37°C. A 10 μl portion was sampled at every 1 minute, and two samples were transferred into a tube in which 20 μl of an MB buffer (40 mM sodium acetate, 100 mM NaCI, 2 mM ZnCl₂, 10% glycerol, pH 4.5) is contained and which is placed on ice. After completion of the sampling, five tubes thus obtained were maintained at 65°C for 10 minutes to inactivate the enzyme, five units of mung bean nuclease were added, and the mixture was maintained at 37°C for 30 minutes. After the reaction, ten DNA fragments different from each other in the degrees of deletion were recovered for each plasmid by agarose gel electrophoresis. The DNA fragments thus recovered were blunt ended with the Klenow fragment, subjected to the ligation reaction at 16°C overnight, and <u>Escherichia coli</u> JM109 was transformed. A single stranded DNA was prepared from each of various clones thus obtained with a helper phage M13K07, and subjected to the sequence reaction with a fluorescent primer cycle-sequence kit available from Applied Biosystem (K.K.), and the DNA sequence was determined with an automatic sequencer.

The DNA sequence comprising 1631 base pairs (bp) thus obtained is illustrated in Figs. 16 - 18 (SEQ ID NO: 7). As a result of examining an open reading frame having a ribosome binding site in front of the initiating codon, two open reading frames which can encode the corresponding proteins (A - B (nucleotide positions 99 - 824 of SEQ ID NO: 7), C - D (nucleotide positions 824 - 1309) in Figs. 16 - 18 were found at the positions where the two xanthophyll synthesis genes crtw and crtw are expected to be present.

Example 12: Identification of pigments produced by Escherichia coli having an Alcaligenes xanthophyll synthesis gene cluster

(1) Identification of astaxanthin and 4-ketozeaxanthin

A deletion plasmid (having only <u>crtW</u>) having a deletion from the right <u>Bst</u>Ell to the nucleotide position 1162 (Fig. 17) (nucleotide position 1162 of SEQ ID NO: 7) among the deletion plasmids from pPC17 prepared in Example 11 was referred to as pPC17-3 (Fig. 19).

The zeaxanthin-producing Escherichia coli JM101 (Example 7 (1)) having pPC17-3 introduced thereinto (Escherichia coli (pACCAR25 Δ crtX, pPC17-3); exhibiting orange) was cultured in 2 liters of 2YT culture medium containing 150 µg/ml of Ap and 30 µg/ml of Cm at 37°C for 18 hours. Bacterial cells collected from the culture solution was extracted with 300 ml of acetone, concentrated, extracted twice with 200 ml of chloroform/methanol (9/1), and concentrated to dryness. Then, thin layer chromatography (TLC) was conducted by dissolving the residue in a small amount of chloroform/methanol (9/1) and developing on a silica gel plate for preparative TLC manufactured by Merck with chloroform/methanol (15/1). The original orange pigment was separated into three spots at the Rf values of 0.54 (ca. 25%),

0.72 (ca. 30%) and 0.91 (ca. 25%). The pigments at the Rf values of 0.54 and 0.72 were scratched off from the TLC plate, dissolved in a small amount of chloroform/methanol (9/1) or methanol, and chromatographed on a Sephadex LH-20 column (15×300 mm) with an eluent of chloroform/methanol (9/1) or methanol to give purified materials in a yield of about 1 mg, respectively.

The materials were identified as 4-ketozeaxanthin (Rf 0.54) and astaxanthin (Rf 0.72), since all of the data of their UV-visible, FD-MS spectra and mobility in TLC (developed with chloroform/methanol (15/1)) accorded with those of the standard samples of 4-ketozeaxanthin and astaxanthin. In addition, the pigment at the Rf value of 0.91 was canthaxanthin (Example 12 (2)).

It was also confirmed by the similar analytical procedures that the β-carotene-producing <u>Escherichia coli</u> JM101 having pPC11 or pPC17 introduced thereinto (<u>Escherichia coli</u> (pACCAR16ΔcrtX, pPC11 or pPC17) (exhibiting orange) produces astaxanthin, 4-ketozeaxanthin and canthaxanthin. Furthermore, it was also confirmed with the authentic sample of phoenicoxanthin obtained in Example 6 that these <u>E. coli</u> transformants produce a trace amount of phoenicoxanthin

(2) Identification of canthaxanthin

The β-carotene-producing Escherichia coli JM101 having pPC17-3 introduced thereinto (Escherichia coli (pACCAR16ΔcrtX, pPC17-3); exhibiting orange) was cultured in 2 liters of 2YT culture medium containing 150 μg/ml of Ap and 30 μg/ml of Cm at 37°C for 18 hours. Bacterial cells collected from the culture solution was extracted with 300 ml of acetone, concentrated, extracted twice with 200 ml of chloroform/methanol (9/1), and concentrated to dryness. Then, thin layer chromatography (TLC) was conducted by dissolving the residue in a small amount of chloroform/methanol (9/1) and developing on a silica gel plate for preparative TLC manufactured by Merck with chloroform/methanol (50/1). The darkest pigment corresponding to 40% of the total amount of orange pigments was scratched off from the TLC plate, dissolved in a small amount of chloroform/methanol (9/1) or chloroform/methanol (1/1), and chromatographed on a Sephadex LH-20 column (15 × 300 mm) with an eluent of chloroform/methanol (9/1) or chloroform/methanol (1/1) to give a purified material in a yield of 2 mg.

The material was identified as canthaxanthin, since all of the data of its UV-visible, FD-MS (m/e 564) spectra and mobility in TLC (developed with chloroform/methanol (50/1)) accorded with those of the standard sample of canthaxanthin (manufactured by BASF). In addition, the pigment of which amount corresponds to 50% of the total amount of the orange pigments observed in the initial extract was considered to be echinenone from its UV-visible spectrum, mobility in silica gel TLC (developed with chloroform/methanol (50/1)), and mobility in HPLC with NOVA PACK HR 6μ C18 (3.9 \times 300 mm; manufactured by Waters) (developed with acetonitrile/methanol/2-propanol (90/6/4)) (Example 7 (2)). In addition, the balance of the extracted pigments, 10%, was unreacted β -carotene.

(3) Identification of zeaxanthin

A plasmid having a 1.15 kb <u>Sal</u>I fragment within pPC11 inserted in the same direction as the plasmid pPC11 into the <u>Sal</u>I site of pBluescript II SK+ was prepared (referred to as pPC13, see Fig. 19).

The β-carotene-producing Escherichia coli JM101 having pPC13 introduced thereinto (Escherichia coli (pACCAR16ΔcrtX, pPC13); exhibiting yellow) was cultured in 2 liters of 2YT culture medium containing 150 μg/ml of Ap and 30 μg/ml of Cm at 37°C for 18 hours. Bacterial cells collected from the culture solution was extracted with 300 ml of acetone, concentrated, extracted twice with 200 ml of chloroform/methanol (9/1), and concentrated to dryness. Then, thin layer chromatography (TLC) was conducted by dissolving the residue in a small amount of chloroform/methanol (9/1) and developing on a silica gel plate for preparative TLC manufactured by Merck with chloroform/methanol (9/1). The darkest pigment corresponding to 90% of the total amount of orange pigments was scratched off from the TLC plate, dissolved in a small amount of chloroform/methanol (9/1) or methanol, and chromatographed on a Sephadex LH-20 column (15 × 300 mm) with an eluent of chloroform/methanol (9/1) or methanol to give a purified material in a yield of 3 mg.

The material was identified as zeaxanthin, since all of the data of its UV-visible, FD-MS (m/e 568) spectra and mobility in TLC (developed with chloroform/methanol (9/1)) accorded with those of the standard sample of zeaxanthin (Example 7 (3)). In addition, the pigment of which amount corresponds to 10% of the total amount of the orange pigments observed in the initial extract was considered to be β -cryptoxanthin from its UV-visible spectrum, mobility in silica gel TLC (developed with chloroform/methanol (9/1)), and mobility in HPLC with NOVA PACK HR $\beta\mu$ C18 (3.9 × 300 mm; manufactured by Waters) (developed with acetonitrile/methanol/2-propanol (90/6/4)) (Example 7 (3)).



Example 13: Identification of the Alcaligenes xanthophyll synthesis gene cluster

(1) Identification of a keto group-introducing enzyme gene

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It is apparent from the results of Examples 11 and 12 (1) that all of the genes required for the synthesis of astaxanthin from β-carotene among the 2.35 kb Pstl fragment contained in pPC11 is contained in the 1.63 kb Pstl - BstEII fragment (pPC17, Fig. 19) in the left side. Thus, the 0.72 kb BstEll - PstI fragment in the right side is not needed. Unique Small and Sall sites are present within the 1.63 kb Pstl - BstEll fragment of pPC17 (Fig. 19). It is confirmed by the pigment analysis with a β-carotene-producing Escherichia coli having the deletion plasmids introduced thereinto that the keto group-introducing enzyme activity was lost when the 0.65 kb and 0.69 kb fragments at the left side from Small and Sall sites were removed. It was also confirmed by the pigment analysis with a β-carotene-producing Escherichia coli having the plasmid introduced thereinto that the plasmid having a 0.69 kb PstI - SalI fragment positioned at the left side of the 1.63 kb Pstl - BstEII fragment inserted into the Pstl - Sall site of pBluescript SK+ has no keto group-introducing enzyme activity. On the other hand, the deletion plasmid pPC17-3 (Fig. 19) in which deletion from the Bst Ell end at the right end to the nucleotide No. 1162 (nucleotide position 1162 in SEQ ID NO: 7) occurred has a keto group-introducing enzyme activity (Example 12 (1), (2)), so that it is considered a gene encoding a keto group-introducing enzyme having an enzyme activity for synthesizing canthaxanthin or astaxanthin with a substrate of β-carotene or zeaxanthin is present in the 1162 bp fragment in pPC17-3, and the aforementioned Small and Sall sites are present within this gene. As a result of determining the nucleotide sequence, an open reading frame which corresponds to the gene and has a ribosome binding site just in front of the initiation codon was successfully detected, so that it was referred to as the crtW gene. The nucleotide sequence of the artw gene and the encoded amino acid sequence are illustrated in Figs. 13 - 14 (SEQ ID NO: 5).

The <u>crtW</u> gene product (CrtW) of <u>Alcaligenes</u> sp. PC-1 has an enzyme activity for converting a methylene group at the 4-position of a β-ionone ring into a keto group, and one of the specific examples is an enzyme activity for synthesizing canthaxanthin from β-carotene as a substrate by way of echinenone (Example 12 (2); see Fig. 11). Furthermore, the <u>crtW</u> gene product also has an enzyme activity for converting a methylene group at the 4-position of a 3-hydroxy-β-ionone ring into a keto group, and one of the specific examples is an enzyme activity for synthesizing astaxanthin from zeaxanthin as a substrate by way of 4-ketozeaxanthin (Example 12 (1); see Fig. 11). In addition, polypeptides having such enzyme activities and DNA strands encoding these polypeptides have not hitherto been known, and the polypeptides and the DNA strands encoding these polypeptides have no total homology to any polypeptides or DNA strands having been hitherto known. Also, the <u>crtW</u> gene products (CrtW) of <u>Agrobacterium aurantiacus</u> sp. nov. MK1 and <u>Alcaligenes</u> sp. PC-1 share high homology (identity of 83%) at the level of amino acid sequence, and the functions of both enzymes are the same. The amino acid sequence in the region of 17% having no identity among these amino acid sequences is considered not so significant to the functions of the enzyme. It is thus considered particularly in this region that a little amount of substitution by the other amino acids, deletion, or addition of the other amino acids will not affect the enzyme activity.

It can be said the keto group-introducing enzyme gene $\underline{\operatorname{crt}W}$ of marine bacteria encodes the β -ionone or 3-hydroxy- β -ionone ring ketolase which converts directly the methylene group at the 4-position into a keto group irrelative to whether a hydroxyl group is added to the 3-position or not. In addition, no such informations have hitherto been described that a methylene group of not only a β -ionone ring and a 3-hydroxy- β -ionone ring but also the other compounds is directly converted into a keto group with one enzyme.

(2) Identification of a hydroxyl group-introducing enzyme gene

All of the genes rerquired for the synthesis of astaxanthin from β -carotene is contained in the 1.63 kb <u>Pst</u>I - <u>Bst</u>Ell fragment (Fig. 19) of pPC17. One <u>Sal</u>I site is present within the 1.63 kb <u>Pst</u>I - <u>Bst</u>Ell fragment of pPC17. It is apparent from the results of Example 12 (3) that a hydroxyl group-introducing enzyme activity is present in a fragment at the right side from the <u>Sal</u>I site. It is thus understood that the hydroxyl group-introducing enzyme activity is present in the 0.94 kb <u>Sal</u>I - <u>Bst</u>Ell fragment which is the right fragment in the 1.63 kb <u>Pst</u>I - <u>Bst</u>Ell fragment. As a result of determining the nucleotide sequence, an open reading frame which corresponds to the gene and has a ribosome binding site just in front of the initiation codon was successfully detected, it was referred to as the <u>crtZ</u> gene. The nucleotide sequence of the <u>crtZ</u> gene and the encoded amino acid sequence are illustrated in Fig. 15 (SEQ ID NO: 6).

The $\underline{\operatorname{crtZ}}$ gene product (CrtZ) of Alcaligenes sp. PC-1 has an enzyme activity for adding a hydroxyl group to the 3-carbon of a β -ionone ring, and one of the specific examples is an enzyme activity for synthesizing zeaxanthin from β -carotene as a substrate by way of β -cryptoxanthin (Example 12 (3); see Fig. 11). Furthermore, the $\underline{\operatorname{crtZ}}$ gene product also has an enzyme activity for adding a hydroxyl group to the 3-carbon of a 4-keto- β -ionone ring, and one of the specific examples is an enzyme activity for synthesizing astaxanthin from canthaxanthin as a substrate by way of phoenicoxanthin (Example 12 (1); see Fig. 11). In addition, polypeptides having the latter enzyme activity and DNA strands encoding these polypeptides have not hitherto been known. Also, the CrtZ of Alcaligenes sp. PC-1 showed significant

homology to the CrtZ of Erwinia uredovora (identity of 58%) at the level of amino acid sequence. In addition, the crtZ gene products (CrtZ) of Agrobacterium aurantiacus sp. nov. MK1 and Alcaligenes sp. PC-1 have high homology (identity of 90%) at the level of amino acid sequence, and the functions of both enzymes are the same. The amino acid sequence in the region of 10% having no identity among these amino acid sequences is considered not so significant to the functions of the enzyme. It is thus considered particularly in this region that a little amount of substitution by the other amino acids, deletion, or addition of the other amino acids will not affect the enzyme activity.

(3) Consideration on minor biosynthetic pathways of xanthophylls

It has been elucidated by our studies with carotenoid synthesis genes of the epiphytic bacterium Erwinia or the photosynthetic bacterium Rhodobacter that carotenoid biosynthesis enzymes generally act by recognizing the half of a carotenoid molecule as a substrate. By way of example, the lycopene cyclase gene of Erwinia, crtY, recognizes the halves of the lycopene molecule to cyclize it. When the phytoene desaturase gene crtl of Rhodobacter was used for the synthesis of neurosporene in place of lycopene in Escherichia coli and crtY of Enwinia was allowed to work on it, the crtY gene product recognizes the half molecular structure common to lycopene to produce a half cyclized β-zeacarotene (Linden, H., Misawa, N., Chamovits, D., Pecher, I., Hirschberg, J., Sandmann, G., "Functional Complementation in Escherichia coli of Different Phytoene Desaturase Genes and Analysis of Accumulated Carotenes", Z. Naturforsch., 46c, p. 1045-1051, 1991). Also, in the present invention, when CrtW is allowed to work on β-carotene or zeaxanthin, echinenone or 4-ketozeaxanthin in which one keto group has been introduced is first synthesized, and when CrtZ is allowed to work on β -carotene or canthaxanthin, β -cryptoxanthin or phoenicoxanthin in which one hydroxyl group has been introduced is first synthesized. It can be considered because these enzymes recognize the half molecule of the substrate. Thus, while Escherichia coli having the crtE, crtB, crtI and crtY genes of Erwinia and the crtZ gene of a marine bacterium produces zeaxanthin as described above, β -cryptoxanthin which is β -carotene having one hydroxyl group introduced thereinto can be detected as an intermediate metabolite. It can be thus considered that if CrtW is present, 3'-hydroxyechinenone or 3-hydroxyechinenone can be synthesized from β-cryptoxanthin as a substrate, and that phoenicoxanthin can be further synthesized by the action of CrtW on these intermediates. The present inventors have not identified these ketocarotenoids in the culture solutions, and the reason is considered to be that only a trace amount of these compounds is present under the conditions carried out in the present experiments. In fact, it was described that 3-hydroxyechinenone or 3'-hydroxyechinenone was detected as a minor intermediate metabolite of astaxanthin in a marine bacterium Agrobacterium aurantiacus sp. nov. MK1 as a gene source (Akihiro Yokoyama ed., "For the biosynthesis of astaxanthin in marine bacteria", Nippon Suisan Gakkai, Spring Symposium, 1994, Abstract, p. 252, 1994). It can be considered from the above descriptions that minor metabolic pathways shown in Fig. 20 are also present in addition to the main metabolic pathways of astaxanthin shown in Fig. 11.

Industrial Applicability

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According to the present invention, the gene clusters required for the biosynthesis of keto group-containing xanthophylls such as astaxanthin, phoenicoxanthin, 4-ketozeaxanthin, canthaxanthin and echinenone have successfully been obtained from marine bacteria, and their structures, nucleotide sequences, and functions have been elucidated. The DNA strands according to the present invention are useful as genes capable of affording the ability of biosynthesis of keto group-containing xanthophylls such as astaxanthin to microorganisms such as Escherichia coli and the like.

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SEQ ID NO: SEQUENCE LENGTH: 639 SEOUENCE TYPE: STRANDEDNESS: double TOPOLOGY: linear 10 MOLECULE TYPE: genomic DNA ORIGINAL SOURCE: ORGANISM: Agrobacterium aurantiacus 15 STRAIN: sp. nov. MK1 SEQUENCE 20 48 GTG CAT GCG CTG TGG TTT CTG GAC GCA GCG GCG CAT CCC ATC CTG GCG Met His Ala Leu Trp Phe Leu Asp Ala Ala Ala His Pro Ile Leu Ala 25 5 10 15 1 96 ATC GCA AAT TTC CTG GGG CTG ACC TGG CTG TCG GTC GGA TTG TTC ATC Ile Ala Asn Phe Leu Gly Leu Thr Trp Leu Ser Val Gly Leu Phe Ile 25 3.0 20 35 ATC GCG CAT GAC GCG ATG CAC GGG TCG GTG GTG CCG GGG CGT CCG CGC 144 Ile Ala His Asp Ala Met His Gly Ser Val Val Pro Gly Arg Pro Arg 40 45 35 40 GCC AAT GCG GCG ATG GGC CAG CTT GTC CTG TGG CTG TAT GCC GGA TTT 192 45 Ala Asn Ala Ala Met Gly Gln Leu Val Leu Trp Leu Tyr Ala Gly Phe 50 55 60 50

SEQUENCE LISTING

	TCG TGG	CGC AA	G ATG AT	C GTC AA	G CAC	ATG GCC	CAT CAC	CGC CAT	GCC	240
5	Ser Trp	Arg Ly	s Met II	e Yal Ly	s His	Met Ala	His His	Arg His	Ala	
	65		. 7	0		75			80	
	GGA ACC	GAC GA	C GAC CC	C GAT TT	C GAC	CAT GGC	GGC CCG	GTC CGC	TGG	288
10	Gl ₃ Thr	Asp As	Asp Pr	o Asp Ph	e Asp	His Gly	Gly Pro	Val Arg	Trp	
			85 .			9 0		95		
15	TAC GCC	CGC TT	C ATC GG	C ACC TA	T TTC	GGC TGG	CGC GAG	GGG CTG	CTG	336
	Tyr Ala	Arg Ph	elle Gl	y Thr Ty	r Phe	Gly Trp	Arg Glu	Gly Leu	Leu	
20		10)		105			110		
	CTG CCC	GTC AT	C GTG AC	G GTC TA	T GCG	CTG ATC	CTT GGG	GAT CGC	TGG	384
	Leu Pro	Yal II	e Val Tb	r Yal Ty	r Ala	Leu Ile	Leu Gly	Asp Arg	Trp	
25		115		1 2	! 0		125			
	ATG TAC	GTG GT	TTC TG	G CCG CT	G CCG	TCG ATC	CTG GCG	TCG ATC	CAG	432
30	Met Tyr	Yal Ya	Phe Tr	p Pro Le	u Pro	Ser Ile	Leu Ala	Ser Ile	Gln	
	130			135			140			
<i>35</i>	CTG TTC	GTG TT	C GGC AC	C TGG CT	G CCG	CAC CGC	CCC GGC	CAC GAC	GCG	480
	Leu Phe	Val Pho	Gly Th	r Trp Le	u Pro	His Arg	Pro Gly	His Asp	Ala	
	145		15			155			160	
40		GAC CG	CAC AA	T GCG CG	G TCG	TCG CGG	ATC AGC	GAC CCC	GTG	528
								Asp Pro		
45		•	165			170		175		
	TCG CTG	CTG ACC	TGC TT	T CAC TT	T GGC	GGT TAT	CAT CAC	GAA CAC	CAC	576
50								Glu His		
		180			185	-		190		

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	CTG	CAC	CCG	ACG	GTG	CCG	TGG	TGG	CGC	CTG	CCC	YCC	ACC	CGC	YCC	AAG	624
5	Leu	His	Pro	Thr	V a I	Pro	Trp	Trp	Arg	Leu	Pro	Ser	Tbr	Arg	Thr	Lys	
			195					200					205				
10	GGG	GAC	ACC	GCA	TGA												639
	Gly	Asp	Thr	Ala	***												
15		210															
20																	

SEQ ID NO: 2 489 SEQUENCE LENGTH: 5 SEQUENCE TYPE: STRANDEDNESS: double TOPOLOGY: linear MOLECULE TYPE: genomic DNA 10. ORIGINAL SOURCE: Agrobacterium aurantiacus ORGANISM: sp. nov. MKl STRAIN: 15 SEQUENCE 20 ATG ACC AAT TTC CTG ATC GTC GCC ACC GTG CTG GTG ATG GAG TTG 48 Met Thr Asn Phe Leu Ile Val Val Ala Thr Val Leu Val Met Glu Leu 25 15 1 5 10 ACG GCC TAT TCC GTC CAC CGC TGG ATC ATG CAC GGC CCC CTG GGC TGG 30 The Ala Tyr Ser Val His Arg Trp Ile Met His Gly Pro Leu Gly Trp 30 25 20 GGC TGG CAC AAG TCC CAC CAC GAG GAA CAC GAC CAC GCG CTG GAA AAG 144 35 Gly Trp His Lys Ser His His Glu Glu His Asp His Ala Leu Glu Lys 40 45 40 35 AAC GAC CTG TAC GGC CTG GTC TTT GCG GTG ATC GCC ACG GTG CTG TTC 192

50 55 60

Asn Asp Leu Tyr Gly Leu Val Phe Ala Val Ile Ala Thr Val Leu Phe

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	ACG	GTG	GGC	TGG	ATC	TGG	GCG	CCG	GTC	CTG	TGG	TGG	ATC	GCC	TTG	GGC	240
5	Thr	Yal	Gly	Trp	I,l e	Trp	Ala	Pro	Y a l	Leu	Trp	Trp	lle	Ala	Leu	Gly	
	65	•				70					75					80	
10	ATG	ACT	GTC	TAT	GGG	CTG	ATC	TAT	TTC	GTC	CTG	CAT	GAC	GGG	CTG	GTG	288
	Met	Thr	Yal	Tyr	Gly	Leu	[] e	1 y T	Phe	V a i	Leu	His	Аsр	Gly	Leu	V a l	
15					85					90					95		
	CAT	CAG	CGC	TGG	CCG	TTC	CGT	TAT	ATC	CCG	CGC	AAG	GGC	TAT	GCC	AGA	336
20	His	Gln	y t g	Trp	Pro	Phe	λrg	Tyr	l l e	Pro	Arg	Lys	Gly	Tyr	Ala	Arg	
			•	100					105					110			
	CGC	CTG	TAT	CAG	GCC	CAC	CGC	CTG	CAC	CAT	GCG	GTC	GAG	GGG	CGC	GAC	384
25	Arg	Leu	Tyr	Gln	Аlа	His	Arg	Leu	His	His	Ala	Y a l	Glu	Gly	Årg	Аsр	
			115					120					125				
30	CAT	TGC	GTC	AGC	TTC	GGC	TTC	ATC	TAT	GCG	CCC	CCG	GTC	GAC	AAG	CTG	432
	His	Суs	Yal	Ser	Phe	Gly	Phe	ll e	Tyr	Ala	Pro	Pro	Val	Asp	Lys	Leu	
35		130					135					140					
	AAG	CAG	GAC	CTG	AAG	ATG	TCG	GGC	GTG	CTG	CGG	GCC	GAG	GCG	CAG	GAG	480
40	Lys	Gln	Asp	Leu	Lys	Met	Ser	Gly	V a l	Lev	Årg	Ala	Glu	Ala	Gln	Glu	٠,
	145					150					155					160	
45	CGC	ACG	TGA														489
	Arg	Thr															
50			* * *														

SEQ ID NO: 3

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SEQUENCE LENGTH: 1161

SEQUENCE TYPE: STRANDEDNESS: double

TOPOLOGY: linear

MOLECULE TYPE: genomic DNA

ORIGINAL SOURCE:

ORGANISM: Agrobacterium aurantiacus

STRAIN: sp. nov. MK1

SEQUENCE

GTG ACC CAT GAC GTG CTG CTG GCA GGG GCG GGC CTT GCC AAC GGG CTG 48

Met Thr His Asp Val Leu Leu Ala Gly Ala Gly Leu Ala Asn Gly Leu

1 5 10 15

ATC GCC CTG GCG CTG CGC GCG GCG CGC CGC GAC CTG CGC GTG CTG CTG 96

lle Ala Leu Ala Leu Arg Ala Ala Arg Pro Asp Leu Arg Val Leu Leu

20 25 30

CTG GAC CAT GCC GCA GGA CCG TCA GAC GGC CAC ACC TGG TCC TGC CAC 144

Leu Asp His Ala Ala Gly Pro Ser Asp Gly His Thr Trp Ser Cys His

35 40 45

GAC CCC GAC CTG TCG CCG GAC TGG CTG GCG CGG CTG AAG CCC CTG CGC 192

Asp Pro Asp Leu Ser Pro Asp Trp Leu Ala Arg Leu Lys Pro Leu Arg

50 55 60

	CGC	GCC	AAC	TGG	CCC	GAC	CAG	GYC	GTG	CGC	TTT	CCC	CGC	CAT	GCC	CGG	240
5	λrg	Ala	A s n	Trp	Pro	Аsр	Gln	Glo	Y a l	λrg	Phe	Pro	γιβ	His	Ala	Åtg	
	65					70					7 5	j				80	
	CGG	CTG	GCC	ACC	GGT	TAC	GGG	TCG	CTG	GAC	GGG	GCG	GCG	CTG	GCG	GAT	288
10	ķιg	Leu	Αla	Thr	Gly	Tyr	Gly	Ser	Leu	Ásp	Gly	Ala	Ala	Leu	Ala	Asp	
					85					90					95		
15 <u>.</u>	GCG	GTG	GTC	CGG	TCG	GGC	GCC	GYC	ATC	CGC	TGG	GAC	AGC	GAC	ATC	GCC	336
	Ala	V a l	Y a l	Arg	Ser	Gly	λla	Glu	l l e	Arg	Trp	Ås p	Ser	Asp	He	Ala	
20				100					105					110			
	CTG	CTG	GAT	GCG	CAG	GGG	GCG	ACG	CTG	TCC	TGC	GGC	ACC	CGG	ATC	GAG	384
25	Leu	Leu	Asp	Ala	Gln	Gly	λla	1 d T	Leu	Ser	Суs	Gly	Thr	y i ë	lle	Glu	
es .			115					120					125				
	GCG	GGC	GCG	GTC	CTG	GAC	GGG	CGG	GGC	GCG	CAG	CCG	TCG	CGG	CAT	CTG	432
30	ķΙz	Gly	Ala	V a l	Leu	λsp	Gly	Arg	Gly	Ala	Gln	Pro	1 s 2	Arg	His	Leu	
		130					135					140					
35	ACC	GTG	GGT	TTC	CAG	**	TTC	GTG	GGT	GTC	GAG	ATC	GYC	ACC	GAC	CGC	480
	Thr	Yal	Gly	Phe	Gln	Lys	Phe	V a l	Gly	Y a l	Glu	He	Glu	Thr	λsp	Åτg	
10	145					150					155					-160	
	CCC	CAC	GGC	GTG	ccc	CGC	CCG	ATG	ATC	ATG	GAC	GCG	ACC	GTC	Y C C	CAG	528
	Pro	His	Gly	Y a l	Pro	y i g	Pro	Met	lle	Met	λsp	Ala	Thr	Val	Thr	Gln	
! 5					165					170					175		
	CAG	GYC	GGG	TAC	CGC	TTC	ATC	TAT	CTG	CTG	CCC	TTC	TCT	CCG	ACG	CGC	576
50	Gln	A s p	Gly	Tyr	Årg	Phe	ΙΙε	Tyr	Leu	Leu	Pro	Phe	Ser	Pro	Thr	ķιg	
				180	•				185					190			

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	ATC	CTG	ATC	GAG	GAC	λCG	CGC	TAT	TCC	GAT	GGC	GGC	GAT	CTG	GAC	GAC	624
5	l l e	Leu	Ιlε	Glu	A s p	Thr	ğı A	Tyr	Ser	Åsр	Gly	Gly	Ásp	Leu	Аsр	λsp	
		•	195					200			•		205				
·	GAC	GCG	CTG	GCG	GCG	GCG	TCC	CAC	GAC	TAT	GCC	CGC	CAG	CAG	GGC	TGG	672
10-	Ásp	Ala	Leu	Ala	Ala	Ala	1 s 2	His	Asp	Tyr	Ala	Arg	Gln	Gin	Gly	Trp	
		210					215					220				· :	
15	ACC	GGG	GCC	GAG	GTC	CGG	CGC	GAA	CGC	GGC	ATC	CTT	CCC	ATC	GCG	CTG	720
										Gly							
20	225			•		230					235					240	
		CAT	GAT	GCG	GCG	GGC	TTC	TGG	GCC	GAT	CAC	GCG	GCG	GGG	CCT	GTT	768
										Asp							
25					245					250					255		
	CCC	GTG	GGA	CTG	CGC	GCG	GGG	TTC	TTT	CAT	CCG	GTC	ACC	GGC	TAT	TCG	816
30																Ser	
	•			260					265					270			
35	сто		: TAT	G00	GCA	CAG	GTO	GCG	GAC	GTG	GTG	GCG	GGT	CTO	TC(CGG	864
																r Gly	
			275					280					285				
40	ccc	; cc			C GAC	GCC	CT(G CGC	: GG(GCC	: ATC	CGC	GA1	AT T	C GC	G ATC	912
																a lle	
45	• •	29			•		29					301					
	GA			G CG	C CG(C GA			r ct	G CG(C CT	177	G AA	C CG	G AT	G CTG	960
50																t Leu	
			5 n.	. 111	o *** (31		•		•	31					320	
	30	J				0 1	•										

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	TTC	CGC	GGC	TGC	GCG	CCC	GAC	CGG	CGC	TAT	ACC	CTG	CTG	CAG	CGG	TTC	1008
5	Phe	Arg	Gly	Cys	Ála	Pro	Хsр	λιg	¥18	Tyr	idT	Leu	Leu	Gln	Årg	Phe	
					325					330					335		
10	TAC	CGC	ATG	CCG	CAT	GGA	CTG	ATC	GAA	CGG	TTC	TAT	GCC	GGC	CGG	CTG	1056
	Tyr	για	He t	Pro	His	Gly	Leu	lle	Glu	λrg	Phe	1 y 1	Ala	Gly	Arg	Leu	
15				340					345					350			
	AGC	GTG	GCG	GAT	CYC	CTG	CGC	ATC	GTG	YCC	GGC	AAG	CCT	CCC	ATT	CCC	1104
20	Ser	Y a l	Alz	Asp	Gln	Leu	Arg	I l e	Yal	Thr	Gly	Lys	Pro	Pro	He	Pro	
			355					360					365				
25	CTT	GGC	ACG	GCC	ATC	CGC	TGC	CTG	CCC	GAA	CGT	CCC	CTG	CTG	AAG	GAA	1152
20	Leu	Gly	Thr	hlа	lle	ķιg	Cys	Leu	019	Glu	Årg	Pro	Leu	Leu	Lys	Glu	
		370					375					380					
30	AAC	GCA	TGA														1161
	Ås n	Ala	* **														
35	385																

SEQ ID NO: 4

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SEQUENCE LENGTH: 2886

SEQUENCE TYPE: STRANDEDNESS: double

TOPOLOGY: linear

MOLECULE TYPE: genomic DNA

ORIGINAL SOURCE:

ORGANISM: Agrobacterium aurantiacus

STRAIN: sp. nov. MK1

SEQUENCE

GGATCCGGCG ACCTTGCGGC GCTGCGCCGC GCGCCTTTGC TGGTGCCTGG GCCGGGTGGC 60
CCTAGGCCGC TGGAACGCCG CGACGCGGCG CGCGGAAACG ACCACGGACC CGGCCCACCG

CAATGGTCGC AAGCAACGGG GATGGAAACC GGCGATGCGG GACTGTAGTC TGCGCGGATC 120
GTTACCAGCG TTCGTTGCCC CTACCTTTGG CCGCTACGCC CTGACATCAG ACGCGCCTAG

GCCGGTCCGG GGGACAAGAT GAGCGCACAT GCCCTGCCCA AGGCAGATCT GACCGCCACC 180
CGGCCAGGCC CCCTGTTCTA CTCGCGTGTA CGGGACGGGT TCCGTCTAGA CTGGCGGTGG

AGCCTGATCG TCTCGGGCGG CATCATCGCC GCTTGGCTGG CCCTGCATGT GCATGCGCTG 240
TCGGACTAGC AGAGCCCGCC GTAGTAGCGG CGAACCGACC GGGACGTACA CGTACGCGAC

TGGTTTCTGG ACGCAGCGGC GCATCCCATC CTGGCGATCG CAAATTTCCT GGGGCTGACC 300
ACCAAAGACC TGCGTCGCCG CGTAGGGTAG GACCGCTAGC GTTTAAAGGA CCCCGACTGG

	TGGCTGTCGG	TCGGATTGTT	CATCATCGCG	CATGACGCGA	TGCACGGGTC	GGTGGTGCCG	360
5	ACCGACAGCC	AGCCTAACAA	GTAGTAGCGC	GTACTGCGCT	ACGTGCCCAG	CCACCACGGC	
10	GGGCGTCCGC	GCGCCAATGC	GGCGATGGGC	CAGCTTGTCC	TGTGGCTGTA	TGCCGGATTT	420
,,,	CCCGCAGGCG	CGCGGTTACG	CCGCTACCCG	GTCGAACAGG	ACACCGACAT	ACGGCCTAAA	
15	TCGTGGCGCA	AGATGATCGT	CAAGCACATG	GCCCATCACC	GCCATGCCGG	AACCGACGAC	480
	AGCACCGCGT	TCTACTAGCA	GTTCGTGTAC	CGGGTAGTGG	CGGTACGGCC	TTGGCTGCTG	
20							
	GACCCCGATT	TCGACCATGG	CGGCCCGGTC	CGCTGGTACG	CCCGCTTCAT	CGGCACCTAT	540
25	CTGGGGCTAA	AGCTGGTACC	GCCGGGCCAG	GCGACCATGC	GGGCGAAGTA	GCCGTGGATA	
30						GCTGATCCTT	600
	AAGCCGACCG	CGCTCCCCGA	CGACGACGGG	CAGTAGCACT	GCCAGATACG	CGACTAGGAA	
35							
						GTCGATCCAG	660
40	CCCCTAGCGA	CCTACATGCA	CCAGAAGACC	GGCGACGGCA	GCTAGGACCG	CAGCTAGGTC	
						CCCGGACCGC	720
45	GACAAGCACA	AGCCGTGGAC	CGACGGCGTG	GCGGGGCCGG	TGCTGCGCAA	GGGCCTGGCG	
							700
50						CTTTCACTTT	
	GTGTTACGCG	CCAGCAGCGC	CTAGTCGCTG	GGGCACAGCG	ACGACTGGA(; GAAAGTGAAA	
55							

	GGCGGTTATC	ATCACGAACA	CCACCTGCAC	CCGACGGTGC	CGTGGTGGCG	CCTGCCCAGC	840
5	CCGCCAATAG	TAGTGCTTGT	GGTGGACGTG	GGCTGCCACG	GCACCACCGC	GGACGGGTCG	
10	ACCCGCACCA	AGGGGGACAC	CGCATGACCA	ATTTCCTGAT	CGTCGTCGCC	ACCGTGCTGG	900
	TGGGCGTGGT	TCCCCCTGTG	GCGTACTGGT	TAAAGGACTA	GCAGCAGCGG	TGGCACGACC	
			•			:	
15	TGATGGAGTT	GACGGCCTAT	TCCGTCCACC	GCTGGATCAT	GCACGGCCCC	CTGGGCTGGG	960
	ACTACCTCAA	CTGCCGGATA	AGGCAGGTGG	CGACCTAGTA	CGTGCCGGGG	GACCCGACCC	
20			•				
	GCTGGCACAA	GTCCCACCAC	GAGGAACACG	ACCACGCGCT	GGAAAAGAAC	GACCTGTACG	1020
25	CGACCGTGTT	CAGGGTGGTG	CTCCTTGTGC	TGGTGCGCGA	CCTTTTCTTG	CTGGACATGC	
30				TGTTCACGGT			1080
	CGGACCAGAA	ACGCCACTAG	CGGTGCCACG	ACAAGTGCCA	CCCGACCTAG	ACCCGCGGCC	
35							
				TCTATGGGCT			1140
۵	AGGACACCAC	CTAGCGGAAC	CCGTACTGAC	AGATACCCGA	CTAGATAAAG	CAGGACGTAC	
40							
				GTTATATCCC		•	
45	TGCCCGACCA	CGTAGTCGCG	ACCGGCAAGG	CAATATAGGG	CGCGTTCCCG	ATACGGTCTG	
50				CGGTCGAGGG			
	CGGACATAGT	CCGGGTGGCG	GACGTGGTAC	GCCAGCTCCC	CGCGCTGGTA	ACGCAGTCGA	

	TCGGCTTCAT	CTATGCGCCC	CCGGTCGACA	AGCTGAAGCA	GGACCTGAAG	ATGTCGGGCG	1320
5	AGCCGAAGTA	GATACGCGGG	GGCCAGCTGT	TCGACTTCGT	CCTGGACTTC	TACAGCCCGC	
	#CC#CCCCC	CCTCCCCTC	GAGCGCACGT	CYCLCYLCYC	стестестес	CAGGGGGGGG	1380
10			CTCGCGTGCA				2000
15							
15	CCTTGCCAAC	GGGCTGATCG	CCCTGGCGCT	GCGCGCGGCG	CGGCCCGACC	TGCGCGTGCT	1440
	GGAACGGTTG	CCCGACTAGC	GGGACCGCGA	CGCGCGCCGC	GCCGGGCTGG	ACGCGCACGA	
20							
	GCTGCTGGAC	CATGCCGCAG	GACCGTCAGA	CGGCCACACC	TGGTCCTGCC	ACGACCCCGA	1500
2 5	CGACGACCTG	GTACGGCGTC	CTGGCAGTCT	GCCGGTGTGG	ACCAGGACGG	TGCTGGGGCT	
30	CCTGTCGCCG	GACTGGCTGG	CGCGGCTGAA	GCCCCTGCGC	CGCGCCAACT	GGCCCGACCA	1560
	GGACAGCGGC	CTGACCGACC	GCGCCGACTT	CGGGGACGCG	GCGCGGTTGA	CCGGGCTGGT	
35							
	GGAGGTGCGC	TTTCCCCGCC	ATGCCEGGCG	GCTGGCCACC	GGTTACGGGT	CGCTGGACGG	1620
40	CCTCCACGCG	AAAGGGGCGG	TACGGGCCGC	CGACCGGTGG	CCAATGCCCA	GCGACCTGCC	
	GGCGGCGCTG	GCGGATGCGG	TGGTCCGGTC	GGGCGCCGAG	ATCCGCTGGG	ACAGCGACAT	1680
4 5	CCGCCGCGAC	CGCCTACGCC	ACCAGGCCAG	CCCGCGGCTC	TAGGCGACCC	TGTCGCTGTA	
	0.000.000.000	C1#CCCC1CC	GGGCGACGCT	CTCCTCCCCC	1000001100	1000000000	1740
50							1170
	GCGGGACGAC	CTACGCGTCC	CCCGCTGCGA	CAGGACGCCG	IUUUUUIAUU	1000000000	
55							

	GGTCCTGGAC	GGGCGGGGCG	CGCAGCCGTC	GCGGCATCTG	ACCGTGGGTT	TCCAGAAATT	1800
5	CCAGGACCTG	ccceccccc	GCGTCGGCAG	CGCCGTAGAC	TGGCACCCAA	AGGTCTTTAA	
	CCTCCCTCTC	CACATCGAGA	CCC7CCCCC	CLICCCCCCC	CCCCGCCCG3	TGATCATGGA	1860
10 .				GGTGCCGCAC			
15	CGCGACCGTC	ACCCAGCAGG	ACGGGTACCG	CTTCATCTAT	CTGCTGCCCT	TCTCTCCGAC	1920
	GCGCTGGCAG	TGGGTCGTCC	TGCCCATGGC	GAAGTAGATA	GACGACGGGA	AGAGAGGCTG	
20			•				
	GCGCATCCTG	ATCGAGGACA	CGCGCTATTC	CGATGGCGGC	GATCTGGACG	ACGACGCGCT	1980
<i>25</i> .	CGCGTAGGAC	TAGCTCCTGT	GCGCGATAAG	GCTACCGCCG	CTAGACCTGC	TGCTGCGCGA	
30	GGCGGCGGCG	TCCCACGACT	ATGCCCGCCA	GCAGGGCTGG	ACCGGGGCCG	AGGTCCGGCG	2040
	CCGCCGCCGC	AGGGTGCTGA	TACGGGCGGT	CGTCCCGACC	TGGCCCCGGC	TCCAGGCCGC	
35	001100000	1.7.0.7.7.0.0.1	T00000T000	0017017000	CCCCCCTTCT	CCCCCCTCI	2100
					•	GGGCCGATCA	2100
40	GCTTGCGCCG	TAGGAAGGGT	AGCGCGACCG	GGTACTACGC	CGCCCGAAGA	CCCGGCTAGT	
	CGCGGCGGG	CCTGTTCCCG	TGGGACTGCG	CGCGGGGTTC	TTTCATCCGG	TCACCGGCTA	2160
45	GCGCCGCCCC	GGACAAGGGC	ACCCTGACGC	GCGCCCCAAG	AAAGTAGGCC	AGTGGCCGAT	
50	TTCGCTGCCC	TATGCGGCAC	AGGTGGCGGA	CGTGGTGGCG	GGTCTGTCCG	GGCCGCCCGG	2220
	AAGCGACGGG	ATACGCCGTG	TCCACCGCCT	GCACCACCGC	CCAGACAGGC	CCGGCGGGCC	
55							

	CACCGACGCG	CTGCGCGGCG	CCATCCGCGA	TTACGCGATC	GACCGGGCGC	900064006	2280
5	GTGGCTGCGC	GACGCGCCGC	GGTAGGCGCT	AATGCGCTAG	CTGGCCCGCG	CGGCGCTGGC	
10	CTTTCTGCGC	CTTTTGAACC	GGATGCTGTT	CCGCGGCTGC	GCGCCCGACC	GGCGCTATAC	2340
10	GAAAGACGCG	GAAAACTTGG	CCTACGACAA	GGCGCCGACG	CGCGGGCTGG	CCGCGATATG	
15	CCTGCTGCAG	CGGTTCTACC	GCATGCCGCA	TGGACTGATC	GAACGGTTCT	ATGCCGGCCG	2400
	GGACGACGTC	GCCAAGATGG	CGTACGGCGT	ACCTGACTAG	CTTGCCAAGA	TACGGCCGGC	
20		•					
	CCTGAGCGTG	GCGGATCAGC	TGCGCATCGT	GACCGCCAAG	CCTCCCATTC	CCCTTGGCAC	2460
25	CGACTCGCAC	CGCCTAGTCG	ACGCGTAGCA	CTGGCCGTTC	GGAGGGTAAG	GGGAACCGTG	
30	GGCCATCCGC	TGCCTGCCCG	AACGTCCCCT	GCTGAAGGAA	AACGCATGAA	CGCCCATTCG	2520
	CCGGTAGGCG	ACGGACGGGC	TTGCAGGGGA	CGACTTCCTT	TTGCGTACTT	GCGGGTAAGC	
35							
	CCCGCGGCCA	AGACCGCCAT	CGTGATCGGC	GCAGGCTTTG	GCGGGCTGGC	CCTGGCCATC	2580
	GGGCGCCGGT	TCTGGCGGTA	GCACTAGCCG	CGTCCGXAAC	CGCCCGACCG	GGACCGGTAG	
40							
	CGCCTGCAGT	CCGCGGGCAT	CGCCACCACC	CTGGTCGAGG	CCCGGGACAA	GCCCGGCGGG	2640
45	GCGGACGTCA	GGCGCCCGTA	GCGGTGGTGG	GACCAGCTCC	GGGCCCTGTT	CGGGCCGCCC	
50	CGCGCCTATG	TCTGGCACGA	TCAGGGCCAT	CTCTTCGACG	CGGGCCCGAC	CGTCATCACC	2700
	GCGCGGATAC	AGACCGTGCT	AGTCCCGGTA	GAGAAGCTGC	GCCCGGGCTG	GCAGTAGTGG	

GACCCCGATG.	CGCTGAAAGA	GCTGTGGGCC	CTGACCGGGC	AGGACATGGC	GCGCGACGTG	2760
CTGGGGCTAC	GCGACTTTCT	CGACACCCGG	GACTGGCCCG	TCCTGTACCG	CGCGCTGCAC	

ACGCTGATGC CGGTCTCGCC CTTCTATCGG CTGATGTGGC CGGGCGGGAA GGTCTTCGAT 2820
TGCGACTACG GCCAGAGCGG GAAGATAGCC GACTACACCG GCCCGCCCTT CCAGAAGCTA

TACGTGAACG AGGCCGATCC AGGGTCTGGG TCTTGCCGTG CCAGGTGAAG CTGTTGCCGT 2880
ATGCACTTGC TCCGGCTAGG TCCCAGACCC AGAACGGCAC GGTCCACTTC GACAACGGCA

GGATCC 2886

CCTAGG

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SEQ ID NO: 5 SEOUENCE LENGTH: 729 5 SEQUENCE TYPE: STRANDEDNESS: double TOPOLOGY: linear MOLECULE TYPE: genomic DNA 10 ORIGINAL SOURCE: Alcaligenes ORGANISM: sp. PC-1 STRAIN: 15 SEQUENCE 20 ATG TCC GGA CGG AAG CCT GGC ACA ACT GGC GAC ACG ATC GTC AAT CTC 48 Met Ser Gly Arg Lys Pro Gly The The Gly Asp The Ile Val Asn Leu 25 5 10 15 1 GGT CTG ACC GCC GCG ATC CTG CTG TGC TGG CTG GTC CTG CAC GCC TTT 96 30 Gly Leu Thr Ala Ala Ile Leu Leu Cys Trp Leu Yal Leu His Ala Phe 20 30 25 ACG CTA TGG TTG CTA GAT GCG GCC GCG CAT CCG CTG CTT GCC GTG CTG 144 35 The Leu Trp Leu Leu Asp Ala Ala Ala His Pro Leu Leu Ala Val Leu 35 · 40 45 40 TGC CTG GCT GGG CTG ACC TGG CTG TCG GTC GGG CTG TTC ATC ATC GCG 192 Cys Leu Ala Gly Leu Thr Trp Leu Ser Val Gly Leu Phe Ile Ile Ala

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														GCC		240
His	Asp	s I A	⅓e t	His	Gly	Ser	Yal	V a l	910	Gly	λrg	Pro	Arg	Аlа	Asn	
. 65					70					75					80	
GCG	GCG	ATC	GGG	CAA	CTG	GCG	CTG	TGG	CTC	TAT	GCG	GGG	TTC	TCG	TGG	288
λla	Ala	i l e	G!y	Gin	Leu	Ala	Leu	Trp	Leu	171	Ala	Gly	Phe	Ser	qıT	
				85					90					95		
CCC	AAG	CTG	ATC	GCC	AAG	CYC	ATG	ACG	CAT	CAC	CGG	CYC	GCC	GGC	ACC	336
Pro	L7s	Lev	lle	Ala	Lys	His	Met	Thr	His	His	γιβ	His	Ala	Gly	Thr	
			100			•		105					110			
GAC	AAC	GAT	CCC	GAT	TTC	GGT	CYC	GGA	GGG	CCC	GTG	CGC	TGG	TAC	GGC	384
Хsр	Asa	Asp	019	k s p	Phe	GI y	His	Gly	Glī	Pro	V a 1	Årg	Trp	Tyr	Gly	
		115					120					125				
AGC	TTC	GTC	TCC	ACC	TAT	TTC	GGC	TGG	CGY	GAG	GGA	CTG	CTG	CTA	CCG.	432
Sei	Phe	Ύ́2 l	Ser	Thr	Tyr	Phe	Gly	T r-p	Arg	Glu	Gly	Leu	Leu	Leu	Pro	
	130					135					140					
GTG	ATC	GTC	A C C	YCC	TAT	GCG	CTG	ATC	CTG	GGC	GAT	CGC	TGG	ATG	TAT	480
V a l	He	Yal	Thr	1 d T	îyı	Ala	Leu	lle	Leu	Gly	λsp	Αrg	Trp	M e t	r v T	
145					150					155					160	
GTC	: ATC	TTC	TGG	CCG	GTC	CCG	GCC	GTT	CTG	GCG	TCG	ATC	CAG	ATT	TTC	528
V a l	lle	Phe	Tip	Pro	V a l	Pro	Ala	V a l	Leu	Ala	s e r	He	Gln	lle	Phe	
				165					170	1				175		
GTO	710	: GGA	АСТ	TGG	CTC	ccc	CYC	CGC	cco	GGA	CA1	GY(CGAT	TTT	CCC	576
Yal	Phe	Gly	The	Trp	Leu	Pro	His	Årg	Pro	Gly	His	. As	a k s	Phe	Pro	
			180					185					190			

20	Årg	Alai	* * *								•						
25	CGC	GCA	TGA														729
	225					230					235					240	
20	Pro	His	Y a l	Pro	Trp	Trp	Årg	Leu	Pro	Å r g	Thr	Åιg	Lys	Thr	Gly	Gly	
	CCG	CAT	GTG	CCG	TGG	TGG	CGC	CTG	CCT	CGT	ACA	CGC	AAG	ACC	GGA	GGC	720
15		210					215					220					
	Leu	Thr	Сīs	Phe	His	Phe	Gly	Gly	Tyr	His	His	Glu	His	His	Leu	His	
10	CTG	ACC	TGC	TTC	CAT	TTC	GGC	GGC	TAT	CAC	CAC	GAA	CAT	CAC	CTG	CAT	672
			195					200					205				
5	As p	y t g	His	A s n	Åla	Arg	Ser	Thr	Gly	Ile	Gly	λsp	Pro	Leu	Ser	Leu	
	GAC	CGG	CYC	AAC	GCG	AGG	TCG	ACC	GGC	ATC	GGC	GYC	CCG	TTG	TCA	CTA	624

SEQ ID NO: 6

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SEQUENCE LENGTH: 489

SEQUENCE TYPE: STRANDEDNESS: double

TOPOLOGY: linear

MOLECULE TYPE: genomic DNA

ORIGINAL SOURCE:

ORGANISM: Alcaligenes

STRAIN: sp. PC-1

SEQUENCE

ATG ACG CAA TTC CTC ATT GTC GTG GCG ACA GTC CTC GTG ATG GAG CTG 48

Met Thr Gln Phe Leu Ile Val Val Ala Thr Val Leu Val Met Glu Leu

5 10 15

ACC GCC TAT TCC GTC CAC CGC TGG ATT ATG CAC GGC CCC CTA GGC TGG 96

The Ala Tyr Ser Val His Arg Trp Ile Met His Gly Pro Leu Gly Trp

20 25 30

GGC TGG CAC AAG TCC CAT CAC GAA GAG CAC GAC CAC GCG TTG GAG AAG 144

Gly Trp His Lys Ser His His Glu Glu His Asp His Alz Leu Glu Lys

35 40 45

AAC GAC CTC TAC GGC GTC GTC TTC GCG GTG CTG GCG ACG ATC CTC TTC 192

Asn Asp Leu Tyr Gly Val Val Phe Ala Val Leu Ala Thr lie Leu Phe

50 55. 60

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	ACC	GTG	GGC	GCC	TAT	TGG	TGG	CCG	GTG	CTG	TGG	TGG	ATC	GCC	CTG	GGC	240
5	Thr	V a l	Gly	Álа	Tyr	Trp	Trp	Pro	Y a l	Leu	Trp	Trp	lle	Αla	Leu	Gly	
	65		•			70					75					80	
10	ATG	ACG	GTC	TAT	GGG	TTG	ATC	TAT	TTC	ATC	CTG	CYC	GAC	GGG	CTT	GTG	288
	Met	1 d T	Yal	Tyr	Gly	Leu	He	Tyr	Phe	11e	Leu	His	Asp	Gly	Leu	V a l	
15					85					90					95	-	
	CAT	CAA	CGC	TGG	CCG	TTT	CGG	TAT	ATT	CCG	CGG	CGG	GGC	TAT	TTC	CGC	336
20	His	Gln	Arg	Trp	Pro	Phe	Arg	Tyr	I I e	P r o	A r g	Å r g	Gly	T y r	Phe	Arg	
20				100					105					110			
	AGG	CTC	TAC	CAA	GCT	CAT	CGC	CTG	CAC	CAC	GCG	GTC	GAG	GGG	CGG	GAC	384
25	Å r g	Leu	Tyr	Gln	λla	His	λιg	Leu	His	His	Ala	Yal	Glu	Gly	y i y	λsp	
			115					120					125				
30	CAC	TGC	GTC	AGC	TTC	GGC	TTC	ATC	TĀT	GCC	CCA	CCC	GTG	GAC	AAG	CTG	432
	His	Суs	Yal	Ser	Phe	Gly	Phe	Ile	Ţyr	Ala	Pro	Pro	V a 1	A s p	Lys	Leu	
<i>35</i>		130					135					140	٠				
	AAG	CAG	GAT	CTG	AAG	CGG	TCG	GGT	GTC	CTG	CGC	ccc	CAG	GAO	GAG	CGT	480
40	Lys	Gln	ı As p	Leu	Lys	Årg	Ser	Gly	Val	Leu	Arg	g Pro	Gli	ı Ası	Glu	Arg	
	145					150					155	5				160	
4 5	CCG	TCC	TGA														489
	Pro	Sei	* * * *	•													

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SEQ ID NO: 7

SEQUENCE LENGTH:

SEQUENCE TYPE: STRANDEDNESS: double

TOPOLOGY: linear

MOLECULE TYPE: genomic DNA

ORIGINAL SOURCE:

ORGANISM: Alcaligenes

STRAIN:

sp. PC-1

SEQUENCE

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CTGCAGGCCG GGCCCGGTGG CCAATGGTCG CAACCGGCAG GACTGGAACA GGACGGCGGG 60 GACGTCCGGC CCGGGCCACC GGTTACCAGC GTTGGCCGTC CTGACCTTGT CCTGCCGCCC

CCGGTCTAGG CTGTCGCCCT ACGCAGCAGG AGTTTCGGAT GTCCGGACGG AAGCCTGGCA 120 GGCCAGATCC GACAGCGGGA TGCGTCGTCC TGAAAGCCTA CAGGCCTGCC TTCGGACCGT

CAACTGGCGA CACGATCGTC AATCTCGGTC TGACCGCCGC GATCCTGCTG TGCTGGCTGG 180 GTTGACCGCT GTGCTAGCAG TTAGAGCCAG ACTGGCGGCG CTAGGACGAC ACGACCGACC

TCCTGCACGC CTTTACGCTA TGGTTGCTAG ATGCGGCCGC GCATCCGCTG CTTGCCGTGC 240 AGGACGTGCG GAAATGCGAT ACCAACGATC TACGCCGGCG CGTAGGCGAC GAACGGCACG

	TGTGCCTGGC	TGGGCTGACC	TGGCTGTCGG	TCGGGCTGTT	CATCATCGCG	CATGACGCAA	300
5	ACACGGACCG	ACCCGACTGG	ACCGACAGCC	AGCCCGACAA	GTAGTAGCGC	GTACTGCGTT	
					•		
	TGCACGGGTC	CGTGGTGCCG	GGGCGGCCGC	GCGCCAATGC	GGCGATCGGG	CAACTGGCGC	360
10	ACGTGCCCAG	GCACCACGGC	cccgccggcg	CGCGGTTACG	CCGCTAGCCC	GTTGACCGCG	
15	TGTGGCTCTA	TGCGGGGTTC	TCGTGGCCCA	AGCTGATCGC	CAAGCACATG	ACCCATCACC	420
	ACACCGAGAT	ACGCCCCAAG	AGCACCGGGT	TCGACTAGCG	GTTCGTGTAC	TGCGTAGTGG	
20							
	GGCACGCCGG	CACCGACAAC	GATCCCGATT	TCGGTCACGG	AGGGCCCGTG	CGCTGGTACG	480
25	CCGTGCGGCC	GTGGCTGTTG	CTAGGGCTAA	AGCCAGTGCC	TCCCGGGCAC	GCGACCATGC	
30	GCAGCTTCGT	CTCCACCTAT	TTCGGCTGGC	GAGAGGGACT	GCTGCTACCG	GTGATCGTCA	540
	CGTCGAAGCA	GAGGTGGATA	AAGCCGACCG	CTCTCCCTGA	CGACGATGGC	CACTAGCAGT	
05	•						
35	CCACCTATGC	GCTGATCCTG	GGCGATCGCT	GGATGTATGT	CATCTTCTGG	CCGGTCCCGG	600
	GGTGGATACG	CGACTAGGAC	CCGCTAGCGA	CCTACATACA	GTAGAAGACC	GGCCAGGGCC	
40							
	CCGTTCTGGC	GTCGATCCAG	ATTTTCGTCT	TCGGAACTTG	GCTGCCCCAC	CGCCCGGGAC	660
45	GGCAAGACCG	CAGCTAGGTC	TAAAAGCAGA	AGCCTTGAAC	CGACGGGGTG	GCGGGCCCTG	
50	ATGACGATTT	TCCCGACCGG	CACAACGCGA	GGTCGACCGG	CATCGGCGAC	CCGTTGTCAC	720
	TACTGCTAAA	AGGGCTGGCC	GTGTTGCGCT	CCAGCTGGCC	GTAGCCGCTG	GGCAACAGTG	

	TACTGACCTG	CTTCCATTTC	GGCGGCTATC	ACCACGAACA	TCACCTGCAT	CCGCATGTGC	780
5	ATGACTGGAC	GAAGGTAAAG	CCGCCGATAG	TGGTGCTTGT	AGTGGACGTA	GGCGTACACG	
	CGTGGTGGCG	CCTGCCTCGT	ACACGCAAGA	CCGGAGGCCG	CGCATGACGC	AATTCCTCAT	840
10	GCACCACCGC	GGACGGAGCA	TGTGCGTTCT	GGCCTCCGGC	GCGTACTGCG	TTAAGGAGTA	
			#0.1#06.1.GOT	0100000TAT	T000T00100	CCTCC1TT1T	0.00
15	•		TGATGGAGCT				900
,	ACAGCACCGC	TGTCAGGAGC	ACTACCTCGA	CTGGCGGATA	AGGCAGGTGG	CGACCTAATA	
20			• .			•	
25	GCYCGGCCCC	CTAGGCTGGG	GCTGGCACAA	GTCCCATCAC	GAAGAGCACG	ACCACGCGTT	960
25	CĠTGCCGGGG	GATCCGACCC	CGACCGTGTT	CAGGGTAGTG	CTTCTCGTGC	TGGTGCGCAA	
30	GGAGAAGAAC	GACCTCTACG	GCGTCGTCTT	CCCCGTCCTC	GCGACGATCC	TCTTCACCGT	1020
	сстсттсттс	· CTGGAGATGC	CGCAGCAGAA	GCGCCACGAC	CGCTGCTAGG	AGAAGTGGCA	
35							
						TCTATGGGTT	1080
40	CCCGCGGATA	ACCACCGGCC	ACGACACCAC	CTAGCGGGAC	CCGTACTGCC	AGATACCCAA	
	GATCTATTTC	ATCCTGCACG	ACGGGCTTGT	GCATCAACGC	TGGCCGTTTC	GGTATATTCC	1140
45	CTAGATAAAG	TAGGACGTGC	TGCCCGAACA	CGTAGTTGCG	ACCGGCAAAG	CCATATAAGG	
50	606066666	TATTTCCGCA	GGCTCTACCA	AGCTCATCGC	CTGCACCACG	CGGTCGAGGG	1200
						GCCAGCTCCC	
55							

	GCGGGACCAC	TGCGTCAGCT	TCGGCTTCAT	CTATGCCCCA	CCCGTGGACA	AGCTGAAGCA	1260
5	CGCCCTGGTG	ACGCAGTCGA	AGCCGAAGTA	GATACGGGGT	GGGCACCTGT	TCGACTTCGT	
	GGATCTGAAG	CGGTCGGGTG	TCCTGCGCCC	CCAGGACGAG	CGTCCGTCGT	GATCTCTGAT	1320
10	CCTAGACTTC	GCCAGCCCAC	AGGACGCGGG	GGTCCTGCTC	GCAGGCAGCA	CTAGAGACTA	
15	CCCGGCGTGG	CCGCATGAAA	TCCGACGTGC	TGCTGGCAGG	GGCCGGCCTT	GCCAACGGAC	1380
	GGGCCGCACC	GGCGTACTTT	AGGCTGCACG	ACGACCGTCC	CCGGCCGGAA	CGGTTGCCTG	
20							
	TGATCGCGCT	CGCGATCCGC	AAGGCGCGGC	CCGACCTTCG	ССТССТССТС	CTGGACCGTG	1440
25				GGCTGGAAGC			
30	CGGCGGGCGC	CTCGGACGGG	CATACTTGGT	CCTGCCACGA	CACCGATTTG	GCGCCGCACT	1500
30				GGACGGTGCT			
35	остоба	CCTGAAGCCG	ATCAGGCGTG	GCGACTGGCC	CGATCAGGAG	GTGCGGTTCC	1560
				CGCTGACCGG			
40	000000000						
	CACACCATTC	OTODOKKOOD	CGGGCCGGAT	ATGGCTCGAT	CGACGGGCGG	GGGCTGATGC	1620
4 5				TACCGAGCTA			
	GICIUGIANO	CGCTTCCCHO					
50	GTGCGGTGAC	ſ					1631
50			•				
	CACGCCACTG	U	•				

Claims

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- A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting a methylene group at the 4-position of a β-ionone ring into a keto group.
- 2. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting the methylene group at the 4-position of the β-ionone ring into a keto group and having an amino acid sequence substantially of amino acid Nos. 1 - 212 which is shown in the SEQ ID NO: 1.
- A DNA strand hybridizing the DNA strand according to claim 2 and having a nucleotide sequence which encodes a polypeptide having an enzyme activity according to claim 2.
 - 4. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting the methylene group at the 4-position of the β-ionone ring into a keto group and having an amino acid sequence substantially of amino acid Nos. 1 - 242 which is shown in the SEQ ID NO: 5.
 - 5. A DNA strand hybridizing the DNA strand according to claim 4 and having a nucleotide sequence which encodes a polypeptide having an enzyme activity according to claim 4.
- 6. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting β-carotene into canthaxanthin by way of echinenone and having an amino acid sequence substantially of amino acid Nos. 1 212 which is shown in the SEQ ID NO: 1.
- 7. A DNA strand hybridizing the DNA strand according to claim 6 and having a nucleotide sequence which encodes a polypeptide having an enzyme activity according to claim 6.
 - 8. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting β-carotene into canthaxanthin by way of echinenone and having an amino acid sequence substantially of amino acid Nos. 1 242 which is shown in the SEQ ID NO: 5.
 - A DNA strand hybridizing the DNA strand according to claim 8 and having a nucleotide sequence which encodes a polypeptide having an enzyme activity according to claim 8.
- 10. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting
 the methylene group at the 4-position of the 3-hydroxy-β-ionone ring into a keto group.
 - 11. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting the methylene group at the 4-position of the 3-hydroxy-β-ionone ring into a keto group and having an amino acid sequence substantially of amino acid Nos. 1 - 212 which is shown in the SEQ ID NO: 1.
- A DNA strand hybridizing the DNA strand according to claim 11 and having a nucleotide sequence which encodes
 a polypeptide having an enzyme activity according to claim 11.
- 13. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting the methylene group at the 4-position of the 3-hydroxy-β-ionone ring into a keto group and having an amino acid sequence substantially of amino acid Nos. 1 242 which is shown in the SEQ ID NO: 5.
 - 14. A DNA strand hybridizing the DNA strand according to claim 13 and having a nucleotide sequence which encodes a polypeptide having an enzyme activity according to claim 13.
 - 15. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting zeaxanthin into astaxanthin by way of 4-ketozeaxanthin and having an amino acid sequence substantially of amino acid Nos. 1 212 which is shown in the SEQ ID NO: 1.
- 55 **16.** A DNA strand hybridizing the DNA strand according to claim 15 and having a nucleotide sequence which encodes a polypeptide having an enzyme activity according to claim 15.

- 17. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting zeaxanthin into astaxanthin by way of 4-ketozeaxanthin and having an amino acid sequence substantially of amino acid Nos. 1 242 which is shown in the SEQ ID NO: 5.
- 18. A DNA strand hybridizing the DNA strand according to claim 17 and having a nucleotide sequence which encodes a polypeptide having an enzyme activity according to claim 17.
 - 19. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for adding a hydroxyl group to the 3-carbon of the 4-keto-β-ionone ring.
 - 20. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for adding a hydroxyl group to position 3-carbon of the 4-keto-β-ionone ring and having an amino acid sequence substantially of amino acid Nos. 1 162 which is shown in the SEQ ID NO: 2.
- 21. A DNA strand hybridizing the DNA strand according to claim 20 and having a nucleotide sequence which encodes a polypeptide having an enzyme activity according to claim 20.
 - 22. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for adding a hydroxyl group to position 3-carbon of the 4-keto-β-ionone ring and having an amino acid sequence substantially of amino acid Nos. 1 162 which is shown in the SEQ ID NO: 6.
 - 23. A DNA strand hybridizing the DNA strand according to claim 22 and having a nucleotide sequence which encodes a polypeptide having an enzyme activity according to claim 22.
- 24. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting canthaxanthin into astaxanthin by way of phoenicoxanthin and having an amino acid sequence substantially of amino acid Nos. 1 162 which is shown in the SEQ ID NO: 2.
- 25. A DNA strand hybridizing the DNA strand according to claim 24 and having a nucleotide sequence which encodesa polypeptide having an enzyme activity according to claim 24.
 - 26. A DNA strand having a nucleotide sequence which encodes a polypeptide having an enzyme activity for converting canthaxanthin into astaxanthin by way of phoenicoxanthin and having an amino acid sequence substantially of amino acid Nos. 1 162 which is shown in the SEQ ID NO: 6.
 - 27. A DNA strand hybridizing the DNA strand according to claim 26 and having a nucleotide sequence which encodes a polypeptide having an enzyme activity according to claim 26.
- 28. A process for producing a xanthophyll comprising introducing the DNA strand according to any one of daims 1 9 into a microorganism having a β-carotene-synthesizing ability, culturing the transformed microorganism in a culture medium, and obtaining canthaxanthin or echinenone from the cultured cells.
 - 29. A process for producing a xanthophyll comprising introducing the DNA strand according to any one of claims 10 -18 into a microorganism having a zeaxanthin-synthesizing ability, culturing the transformed microorganism in a culture medium, and obtaining astaxanthin or 4-ketozeaxanthin from the cultured cells.
 - 30. A process for producing a xanthophyll comprising introducing the DNA strand according to any one of claims 19 27 into a microorganism having a canthaxanthin-synthesizing ability, culturing the transformed microorganism in a culture medium, and obtaining astaxanthin or phoenicoxanthin from the cultured cells.
 - 31. A process for producing a xanthophyll according to any one of claims 28 30, wherein the microorganism is a bacterium or yeast.

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A GTG CAT GCG CTG TGG TTT CTG GAC GCA GCG GCG CAT CCC ATC CTG GCG ATC GCA Met His Ala Leu Trp Phe Leu Asp Ala Ala Ala His Pro Ile Leu Ala Ile Ala AAT TIC CIG GGG CTG ACC TGG CTG TCG GTC GGA TTG TTC ATC ATC GCG CAT GAC Asn Phe Leu Gly Leu Thr Trp Leu Ser Val Gly Leu Phe Ile Ile Ala His Asp GCG ATG CAC GGG TCG GTG GTG CCG GGG CGT CCG CGC GCC AAT GCG GCG ATG GGC Ala Met His Gly Ser Val Val Pro Gly Arg Pro Arg Ala Asn Ala Met Gly CAG CTT GTC CTG TGG CTG TAT GCC GGA TTT TCG TGG CGC AAG ATG ATC GTC AAG Gln Leu Val Leu Trp Leu Tyr Ala Gly Phe Ser Trp Arg Lys Met Ile Val Lys CAC ATG GCC CAT CAC CGC CAT GCC GGA ACC GAC GAC GAC CCC GAT TTC GAC CAT His Met Ala His His Arg His Ala Gly Thr Asp Asp Pro Asp Phe Asp His GGC GGC CCG GTC CGC TGG TAC GCC CGC TTC ATC GGC ACC TAT TTC GGC TGG CGC Gly Gly Pro Val Arg Trp Tyr Ala Arg Phe Ile Gly Thr Tyr Phe Gly Trp Arg GAG GGG CTG CTG CCC GTC ATC GTG ACG GTC TAT GCG CTG ATC CTT GGG GAT Glu Gly Leu Leu Pro Val Ile Val Thr Val Tyr Ala Leu Ile Leu Gly Asp CGC TGG ATG TAC GTG GTC TTC TGG CCG CTG CCG TCG ATC CTG GCG TCG ATC CAG Arg Trp Met Tyr Val Val Phe Trp Pro Leu Pro Ser Ile Leu Ala Ser Ile Gln CTG TTC GTG TTC GGC ACC TGG CTG CCG CAC CGC CCC GGC CAC GAC GCG TTC CCG Leu Phe Val Phe Gly Thr Trp Leu Pro His Arg Pro Gly His Asp Ala Phe Pro GAC CGC CAC AAT GCG CGG TCG TCG CGG ATC AGC GAC CCC GTG TCG CTG ACC Asp Arg His Asn Ala Arg Ser Ser Arg Ile Ser Asp Pro Val Ser Leu Leu Thr TGC TTT CAC TTT GGC GGT TAT CAT CAC GAA CAC CAC CTG CAC CCG ACG GTG CCG Cys Phe His Phe Gly Gly Tyr His His Glu His His Leu His Pro Thr Val Pro TGG TGG CGC CTG CCC AGC ACC CGC ACC AAG GGG GAC ACC GCA TGA Trp Trp Arg Leu Pro Ser Thr Arg Thr Lys Gly Asp Thr Ala ***

FIG. I

ATG ACC AAT TTC CTG ATC GTC GTC GCC ACC GTG CTG GTG ATG GAG TTG ACG GCC Met Thr Asn Phe Leu Ile Val Val Ala Thr Val Leu Val Met Glu Leu Thr Ala TAT TCC GTC CAC CGC TGG ATC ATG CAC GGC CCC CTG GGC TGG GGC TGG CAC AAG Tyr Ser Val His Arg Trp Ile Met His Gly Pro Leu Gly Trp Gly Trp His Lys TCC CAC GAG GAA CAC GAC CAC GCG CTG GAA AAG AAC GAC CTG TAC GGC CTG Ser His His Glu Glu His Asp His Ala Leu Glu Lys Asn Asp Leu Tyr Gly Leu GTC TTT GCG GTG ATC GCC ACG GTG CTG TTC ACG GTG GGC TGG ATC TGG GCG CCG Val Phe Ala Val Ile Ala Thr Val Leu Phe Thr Val Gly Trp Ile Trp Ala Pro GTC CTG TGG TGG ATC GCC TTG GGC ATG ACT GTC TAT GGG CTG ATC TAT TTC GTC Val Leu Trp Trp Ile Ala Leu Gly Met Thr Val Tyr Gly Leu Ile Tyr Phe Val CTG CAT GAC GGG CTG GTG CAT CAG CGC TGG CCG TTC CGT TAT ATC CCG CGC AAG Leu His Asp Gly Leu Val His Gln Arg Trp Pro Phe Arg Tyr Ile Pro Arg Lys GGC TAT GCC AGA CGC CTG TAT CAG GCC CAC CGC CTG CAC CAT GCG GTC GAG GGG Gly Tyr Ala Arg Arg Leu Tyr Gln Ala His Arg Leu His His Ala Val Glu Gly CGC GAC CAT TGC GTC AGC TTC GGC TTC ATC TAT GCG CCC CCG GTC GAC AAG CTG Arg Asp His Cys Val Ser Phe Gly Phe Ile Tyr Ala Pro Pro Val Asp Lys Leu AAG CAG GAC CTG AAG ATG TCG GGC GTG CTG CGG GCC GAG GCG CAG GAG CGC ACG Lys Gln Asp Leu Lys Met Ser Gly Val Leu Arg Ala Glu Ala Gln Glu Arg Thr TGA

FIG. 2

E 1366 1375 GTG ACC CAT GAC GTG CTG GCA GGG GCG GGC CTT GCC AAC GGG CTG ATC GCC Met Thr His Asp Val Leu Leu Ala Gly Ala Gly Leu Ala Asn Gly Leu Ile Ala CTG GCG CTG CGC GCG CGG CCC GAC CTG CGC GTG CTG CTG GAC CAT GCC Leu Ala Leu Arg Ala Ala Arg Pro Asp Leu Arg Val Leu Leu Asp His Ala GCA GGA CCG TCA GAC GGC CAC ACC TGG TCC TGC CAC GAC CCC GAC CTG TCG CCG Ala Gly Pro Ser Asp Gly His Thr Trp Ser Cys His Asp Pro Asp Leu Ser Pro GAC TGG CTG GCG CGG CTG AAG CCC CTG CGC CGC GCC AAC TGG CCC GAC CAG GAG Asp Trp Leu Ala Arg Leu Lys Pro Leu Arg Arg Ala Asn Trp Pro Asp Gln Glu GTG CGC TTT CCC CGC CAT GCC CGG CGG CTG GCC ACC GGT TAC GGG TCG CTG GAC Val Arg Phe Pro Arg His Ala Arg Arg Leu Ala Thr Gly Tyr Gly Ser Leu Asp GGG GCG GCG GTG GCG GTG GTC CGG TCG GGC GCC GAG ATC CGC TGG GAC Gly Ala Ala Leu Ala Asp Ala Val Val Arg Ser Gly Ala Glu Ile Arg Trp Asp AGC GAC ATC GCC CTG CTG GAT GCG CAG GGG GCG ACG CTG TCC TGC GGC ACC CGG Ser Asp Ile Ala Leu Leu Asp Ala Gln Gly Ala Thr Leu Ser Cys Gly Thr Arg ATC GAG GCG GGC GCG GTC CTG GAC GGG CGG GGC GCG CAG CCG TCG CGG CAT CTG Ile Glu Ala Gly Ala Val Leu Asp Gly Arg Gly Ala Gln Pro Ser Arg His Leu ACC GTG GGT TTC CAG AAA TTC GTG GGT GTC GAG ATC GAG ACC GAC CGC CCC CAC Thr Val Gly Phe Gln Lys Phe Val Gly Val Glu Ile Glu Thr Asp Arg Pro His GGC GTG CCC CGC CCG ATG ATC ATG GAC GCG ACC GTC ACC CAG CAG GAC GGG TAC Gly Val Pro Arg Pro Met Ile Met Asp Ala Thr Val Thr Gln Gln Asp Gly Tyr CGC TTC ATC TAT CTG CTG CCC TTC TCT CCG ACG CGC ATC CTG ATC GAG GAC ACG Arg Phe Ile Tyr Leu Leu Pro Phe Ser Pro Thr Arg Ile Leu Ile Glu Asp Thr CGC TAT TCC GAT GGC GGC GAT CTG GAC GAC GCG CTG GCG GCG GCG TCC CAC Arg Tyr Ser Asp Gly Gly Asp Leu Asp Asp Asp Ala Leu Ala Ala Ser His

GAC TAT GCC CGC CAG GGC TGG ACC GGG GCC GAG GTC CGG CGC GAA CGC GGC Asp Tyr Ala Arg Gln Gln Gly Trp Thr Gly Ala Glu Val Arg Arg Glu Arg Gly ATC CTT CCC ATC GCG CTG GCC CAT GAT GCG GCG GGC TTC TGG GCC GAT CAC GCG Ile Leu Pro Ile Ala Leu Ala His Asp Ala Ala Gly Phe Trp Ala Asp His Ala GCG GGG CCT GTT CCC GTG GGA CTG CGC GCG GGG TTC TTT CAT CCG GTC ACC GGC Ala Gly Pro Val Pro Val Gly Leu Arg Ala Gly Phe Phe His Pro Val Thr Gly TAT TCG CTG CCC TAT GCG GCA CAG GTG GCG GAC GTG GCG GGT CTG TCC GGG Tyr Ser Leu Pro Tyr Ala Ala Gln Val Ala Asp Val Val Ala Gly Leu Ser Gly CCG CCC GGC ACC GAC GCG CTG CGC GGC GCC ATC CGC GAT TAC GCG ATC GAC CGG Pro Pro Gly Thr Asp Ala Leu Arg Gly Ala Ile Arg Asp Tyr Ala Ile Asp Arg GCG CGC CGC GAC CGC TTT CTG CGC CTT TTG AAC CGG ATG CTG TTC CGC GGC TGC Ala Arg Arg Asp Arg Phe Leu Arg Leu Leu Asn Arg Met Leu Phe Arg Gly Cys GCG CCC GAC CGG CGC TAT ACC CTG CTG CAG CGG TTC TAC CGC ATG CCG CAT GGA Ala Pro Asp Arg Arg Tyr Thr Leu Leu Gln Arg Phe Tyr Arg Met Pro His Gly CTG ATC GAA CGG TTC TAT GCC GGC CGG CTG AGC GTG GCG GAT CAG CTG CGC ATC Leu Ile.Glu Arg Phe Tyr Ala Gly Arg Leu Ser Val Ala Asp Gln Leu Arg Ile GTG ACC GGC AAG CCT CCC ATT CCC CTT GGC ACG GCC ATC CGC TGC CTG CCC GAA Val Thr Gly Lys Pro Pro Ile Pro Leu Gly Thr Ala Ile Arg Cys Leu Pro Glu CGT CCC CTG CTG AAG GAA AAC GCA TGA Arg Pro Leu Leu Lys Glu Asn Ala ***

F I G. 4

	10		20		30		40		50		60
GGATC	cecce	acctt tggaa	CGCCG	GCTGC C	SCCGC (cecee :	TTTGC AAACG	TGGTG (CCTGG (GGACC (ceece e	TGGC ACCG
	70		80		90		100		110		120
CAATG	GTCĠC	ARGCA	ACGGG	GATGG A	AAACC	GGCGA	TGCGG	GACTG	TAGTC	TGCGC (GATC
GTTAC	CAGCG	TTCGT	TGCCC	CTACC :	TTTGG	CCGCT	ACGCC	CTGAC	ATCAG	ACGCG (CTAG
	130		140		150		160		170		180
GCCGG	TCCGG	GGGAC	AAGAT	GAGCG (CACAT	GCCCT	GCCCA	AGGCA	GATCT	GACCG (CCACC
CGGCC	AGGCC	CCCTG	TTCTA	CTCGC (GTGTA	CGGGA	CGGGT	TCCGT	CTAGA	CTGGC	GGTGG
	190		200		210		220	•	A 230		240
AGCCT	GATCG	TCTCG	GGCGG	CATCA	TCGCC	GCTTG	GCTGG	CCCTG	CATGT	GCATG	CGCTG
TCGGA	CTAGC	AGAGC	CCGCC	GTAGT	AGCGG	CGAAC	CGACC	GGGAC	GTACA	CGTAC	GCGAC
•	250		260		270		280		290		300
тсстт	TOTEG	ACGCA	GCGGC	GCATC	CCATC	CTGGC	GATCG	CAAAT	TTCCT	GGGGC	TGACC
ACCAA	AGACC	TGCGT	CGCCG	CGTAG	GGTAG	GACCG	CTAGC	GTTTA	AAGGA	CCCCG	ACTGG
,	310		320		330		340		350		360 *
										GGTGG	
ACCGA	CAGCC	AGCCT	AACAA	GTAGT	AGCGC	GTACT	GCGCT	ACGTG	CCCAG	CCACC	ACGGC
	370)	380		390		400		410		420
GGGCG	TCCGC	GCGCC	AATGC	GGCGA	TGGGC	CAGCT	TGTCC	TGTGG	CTGTA	TGCCG	GATTT
										ACGGC	
	430) -	440		450)	460)	470	1	480
TCGT	GCGC	A AGATO	ATCGT	CAAGC	ACATO	GCCCA	TCAC	CGCCAT	GCCGG	AACCG	ACGAC
AGCAC	CGCG	TCTAC	TAGCA	GTTCG	TGTAC	CGGGI	AGTG(G CGGTA	CGGCC	TTGGC	TGCTG
	49	0	500)	510	•	52	D *	530) ·	540
										r CGGCA	
CTGG	GCTA	A AGCTO	G GTACC	c eccee	GCCAG	G GCGAG	CATG	c GGGC	AAGT	A GCCGT	GGATA
	55	0	560	•	57	0 -	58	0	59	D -	600
										C GCTG?	
AAGC	C GACC	G CGCT	c cccg	A CGACG	ACGG	G CAGT	A GCAC	T GCCA	G ATAC	G CGAC1	AGGAA
	61	0	62	0	63	0	6 4	0	65	0	660
acco	1 TOO	T GGET	G TACE	• ה הפירים	፣ ምርጥር	* G ((GC	ጥ ፍርርር	т сеэт •	ר רזינים	* C GTCG	* TCCAG
										G CAGC	

	670		680		690		700		710		720
CTGTT GACAA											
	730		740		750		760		770		780
CACAA GTGTT								TGCTG .			
	790		800		810		820		830		840
							-	CGTGG GCACC			
	850		860	Ć'	870		880		890		900
								CGTCG GCAGC			
10000	910	10000	920	E GCGT		IAAAG	940	GCAGC	950	10001	960
TGATG	GAGTT	GACGG	CCTAT		•	GCTGG	ATCAT	GCACG	GCCCC	CTGGG	* CTGGG
ACTAC	CTCAA	CTGCC	GGATA	AGGCA	GGTGG	CGACC	TAGTA	CGTGC	CGGGG	GACCC	GACCC
	970		980		990		1000		1010		1020
								GGAAA CCTTT			GTACG
•	1030		1040		1050		1060		1070		1080
GCCTG	GTCTT	TGCGG	TGATC	GCCAC	* CGTGC	TGTTC	* CGGT	GGGCT	* GGATC	тесес	sccee
								CCCGA			
	1090		1100		1110		1120		1130		1140
								GATCT CTAGA			
	1150		1160						1190		1200
	*		*		1170		1180		*		*
TGCCC	GACCA	CGTAG	AGCGC	TGGCC	GTTCC CAAGG	GTTAT CAATA	ATCCC TAGGG	GCGCA CGCGT	AGGGC TCCCG	TATGC ATACG	CAGAC GTCTG
	1210		1220		1230		1240		1250		1260
GCCTG	TATCA	GGCCC	ACCGC	CTGCA	CCATO	CGGTC	GAGGG	GCGCG	ACCAT	TGCGT	CAGCT
CGGAC	ATAGT	CCGGG	TGGCG	GACGI	GGTAC	GCCAG	CTCCC	CGCGC	TGGTA	ACGCA	GTCGA
	1270	1	1280	I	1290)	1300		1310)	1320
TCGGC AGCCG	TTCAT AAGTA	CTATG	CGCCC	CCGG1	CGACA	AGCTO	AAGCA	GGACC	TGAAC	TACAG	GGGCG GCCGC

FIG. 6

1330	0	1340	E:	1350 W-		1360		1370		1380
TGCTG CGGGG	- C CGAGG (G GCTCC (CGCAG G	AGCG CA	ACGT G	ACCC A	ATGAC (GTGCT G CACGA C	GACC (CAGGG (STCCC (GCGGG GCCC
139		1400		1410D		1420		1430		1440
CCTTG CCAA GGAAC GGTT	C GGGCT G	GATCG (CCTG G	CGCT G GCGA C	GCGC	- - - - - - - - - - - - - - - - - - -	ccccc (CGACC	TGCGC (ACGCG (STGCT CACGA
145	0	1460		1470		1480		1490		1500
GCTGC TGGA	• C CATGC (G GTACG (+ CGCAG G GCGTC C	SACCG TO	CAGA C GTCT G	GGCC GCCGG	ACACC TGTGG	TGGTC (ACCAG (CTGCC	ACGAC TGCTG	CCCGA GGGCT
151	0	1520		1530		1540		1550		1560
CCTGT CGCC	G GACTG	GCTGG (CGACC (cocc c	TGAA C	ceeee	TGCGC ACGCG	CGCGC (CAACT GTTGA	GGČCC CCGGG	GACCA CTGGT
157	0	1580		1590		1600		1610		1620
GGAGG TGCC	GC TTTCC	ccccc :	ATGCC C	CCCC (GCTGG CGACC	CCACC GGTGG	GGTTA CCAAT	CGGGT GCCCA	CGCTG GCGAC	GACGG CTGCC
163	30	1640		1650		1660		1670 *		1680
cccc ccc	rg gcgga ac cgcct	TGCGG ACGCC	TGGTC (CGGTC	ccccc ccccc	CCGAG	ATCCG TAGGC	CTGGG GACCC	ACAGC TGTCG	GACAT CTGTA
1 5	90	1700		1710		1720		1730		1740
CGCCC TGC	TG GATGC AC CTACG	GCAGG CGTCC	ccccc	ACGCT TGCGA	GTCCT CAGGA	ccccc	ACCCG TGGGC	GATCG CTAGC	AGGCG TCCGC	CCCCC
17	50	1760		1770		1780	!	1790		1800
GGTCC TGG CCAGG ACC	AC GGGCG TG CCCGC	GGGCG	CGCAG GCGTC	CCGTC GGCAG	CCCCC	TAGAC	ACCGT TGGCA	GGGTT CCCA	TCCAG AGGTC	TTAAA AATTT
18	10	1820		1830		1840	•	1850		1860
CGTGG GTG GCACC CAC	TC GAGAT	CGAGA GCTCT	CCGAC GGCTG	CGCCC GCGGG	CCACO	GCGT(cccc c ggggc	GGGC,	A TGATO	ATGGA TACCT
18	370	1980		1890		190	o -	191	0	1920
CGCGA CCC GCGCT GGC	TC ACCCA	A GCAGG CGTCC	ACGGG TGCCC	TACCG	CTTC.	A TCTA T AGAT	T CTGCT A GACGA	GCCC A CGGG	T TCTC	T CCGAC A GGCTG
-1	930	1940)	1950)	196	0	197	0	1980
GCGCA TC	- CTG ATCGA GAC TAGCI	- A GGACA T CCTGT	cGCGC	OTTAT DAATA	CGAT	c cccc	G CTAG	T GGAC	G ACGA C TGCT	C GCGCT G CGCGA

F1G. 7

		1990		2000	•	2010		2020		2030		2040
									ACCGG			
		2050		2060		2070		2080		2090		2100
_		*		*		*		*		*		* C\#C\
									CGCCC			
		2110		2120		2130 -		2140		2150		2160
C	GCGG	CGGGG	CCTGT	TCCCG	TGGGA	CTGCG	CGCGG	GGTTC	TTTCA	TCCGG	TCACC	GGCTA
G	CGCC	GCCCC	GGACA	AGGGC	ACCCT	GACGC	GCGCC	CCAAG	AAAGT	AGGCC	AGTGG	CCGAT
		2170		2180		2190		2200		2210		2220
									GGTCT CCAGA			
		2230		2240		2250		2260		2270		2280
_	*CCG	ACGCG	CTGCG	CGGCG	CCATC		TTACC	•	GACCG		GCCGC	G2CCG
									CTGGC			
		2290		2300		2310		2320		2330		2340
C	TTTC	TGCGC	CTTTT	GAACC	GGATG	CTGTT	CCGCG	GCTGC	GCGCC	CGACC	GGCGC	TATAC
G	AAAG	ACGCG	GAAAA	CTTGG	CCTAC	GACAA	GGCGC	CGACG	CGCGG	GCTGG	CCGCG	ATATG
		2350		2360 *		2370		2380		2390		2400
C	CTGC	TGCAG	CGGTT	CTACC	GCATG	CCGCA	TGGAC	TGATC	GAACG	GTTCT	ATGCC	GGCCG
C	GACG	ACGTC	GCCAA	GATGG	CGTAC	GGCGT	ACCTG	ACTAG	CTTGC	CAAGA	TACGG	CCGGC
		2410		2420		2430		2440)	2450	1	2460
									CCTCC			
(CGACT			AGTCG	ACGCG	TAGCA	CTGGC	CGTTC	GGAGG	GTAAG	GGGAA	CCGTG
		2470		2480		2490	1	2500		2510)	2520
(GGCCA	TCCGC	TGCCT	GCCCG	AACGT	רכככיי	ССТС		A AACGO	. אבפא	הפררנ	TTTCG
									TTGCG			
		2530)	2540		2550		2560	o *	F257	•	2580
	CCCG	GGCC	AGACO	GCCAT	CGTGA	TCGG	GCAG	G CTTTC	G GCGGC	CTGG	CCTG	CCATC
ı	GGGCC	G CCGG1	TCTGG	CGGTA	GCACT	AGCC	CGTC	GAAA	CGCCC	GACC	G GGAC	GGTAG
		2590)	2600)	2610		262	0	263	0	2640
							•		*		*	
	GCGG:	A CGTCA	GCCCC	CCGTA	GCGC3	GGTG	CTGG: GACC	r CGAG(A GCTC(G CCCGC	G GACA C CTGT	A GCCC(T CGGG(ccccc ccccc

	2650		2660		2670		2680		2690		2700
CGCGC	CTATG GATAC	TCTGG AGACC	CACGA GTGCT	TCAGG AGTCC	GCCAT CGGTA	CTCTT GAGAA	ODKDO COTGC	CGGGC	CCGAC GGCTG	CGTCA GCAGT	TCACC AGTGG
	2710		2720		2730		2740		2750		2760
GACCC CTGGG	CGATG GCTAC	CGCTG GCGAC	AAAGA TTTCT	GCTGT CGACA	ccccc	CTGAC GACTG	CGGGC	AGGAC TCCTG	ATGGC TACCG	CCCCC	ACGTG TGCAC
	2770		2780 *		2790 *	•	2800		2610	•	2820
ACGCT TGCGA	GATGC CTACG	CGGTC	TCGCC AGCGG	CTTCT GAAGA	ATCGG TAGCC	CTGAT	CACCG	GCCCG	GGGAA CCCTT	GGTCT	TCGAT AGCTA
	2830		2840		2850		2860 *		2870		2860
TACGT ATGCA	GAACG CTTGC	AGGCC	GATCC CTAGG	AGGGT	CTGGG	TCTTG : AGAAC	CCGTG GGCAC	CCAGG GGTCC	TGAAG ACTTC	GACAA	GCCGT
28	86										
GGATC CCTAG	-			•							

F I G. 11

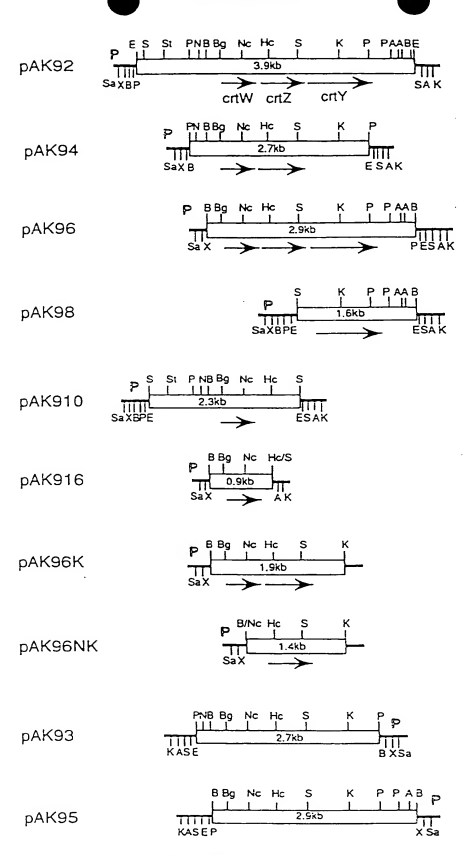


FIG. 12

760 770 780 790 800 810
CATCACCTGCATCCGCATGTGCCGTGGTGGCGCCTTGCTCGTACACGCAAGACCGGAGGC
221 HisHisLeuHisProHisValProTrpTrpArgLeuProArgThrArgLysThrGlyGly

820 827 CGCGCATGA 241 ArgAla***

	10		20		30		40		50		60
										GGACG	
GACGT	CCGGC	CCGGG	CCACC	GGTTA	CCAGC	GTTGG	CCGTC	CTGAC	CTTGT	CCTGC	CGCCC
							Αl				
	70		80		90		Y		110		120
										AAGCC	
GGCCA	GATCC	GACAG	CGGGA	TGCGT	CGTCC	TCAAA	GCCTA	CAGGC	CTGCC	TTCGG	ACCGT
	120		3.40		150		160		170		180
C)) CM	130	CACCA	140	> > TCT		TCACC		GATCC		TGCTG	
										ACGAC	
GIIGA	CCGCI	GIGCI	AGCAG	IIAGA	GCCAG	AC100	00000	C11.00			001.00
	190		200		210		220		230		240
TCCTG		СТТТА		TGGTT		ATGCG		GCATC	CGCTG	CTTGC	CGTGC
										GAACG	
	0.000	0									
	250		260		270		280		290		300
TGTGC	CTGGC	TGGGC	TGACC	TGGCT	GTCGG	TCGGG	CTGTT	CATCA	TCGCG	CATGA	CGCAA
ACACG	GACCG	ACCCG	ACTGG	ACCGA	CAGCC	AGCCC	GACAA	GTAGT	AGCGC	GTACT	GCGTT
	310		320		330		340		350		360
										CAACT	
ACGTG	CCCAG	GCACC	ACGGC	CCCGC	CGGCG	CGCGG	TTACG	CCGCT	AGCCC	GTTGA	CCGCG
									43.0		420
	370		380		390		400		410	10001	420
										ACGCA	
ACACC	GAGAT	ACGCC	CCAAG	AGCAC	CGGGT	TCGAC	INGCG	GIICG	IGIAC	TGCGT	AGIGG
	430		440		450		460		470		480
CCCAC		CACCG		GATCC						CGCTG	
CCGTG	CCCCC	GTGGC	тсттс	CTAGG	GCTAA	AGCCA	GTGCC	TCCCG	GGCAC	GCGAC	CATGC
00010	00000	01000	.0.10	0100							
	490		500		510		520		530		540
GCAGC	TTCGT	CTCCA	CCTAT	TTCGG	CTGGC	GAGAG	GGACT	GCTGC	TACCG	GTGAT	CGTCA
										CACTA	
	550		560		570		580		590		600
										CCGGI	
GGTGG	ATACG	CGACT	AGGAC	CCGCT	AGCGA	CCTAC	ATACA	GTAGA	AGACC	GGCCA	GGGCC
	610		620		630		640		650		660
										CGCCC	
GGCAA	GACCG	CAGCT	AGGTC	AAAA	GCAGA	AGCCT	TGAAC	CGACG	GGGTG	GCGGG	CCCTG
	630						200		716	,	720
> mC > C	670		680		690		700		710		720
										CCGTT GGCA	
TACTG	CTAAA	AGGGC	TOCCC	GTGTT	CCCC	CCAGC		, GINGC		, GGCAA	CVOIG

FIG. 16

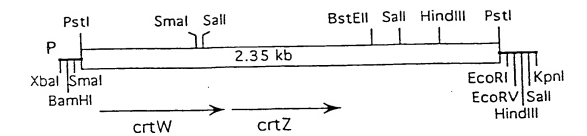
730 740 750 760 770 TACTG ACCTG CTTCC ATTTC GGCGG CTATC ACCAC GAACA TCACC TGCAT CCGCA ATGAC TGGAC GAAGG TAAAG CCGCC GATAG TGGTG CTTGT AGTGG ACGTA GGCGT 790 800 810 820 830 CGTGG TGGCG CCTGC CTCGT ACACG CAAGA CCGGA GGCCG CGCAT GACGC AATTC	780 IGTGC ACACG 840
ATGAC TGGAC GAAGG TAAAG CCGCC GATAG TGGTG CTTGT AGTGG ACGTA GGCGT	ACACG
790 800 810 820 C	
7,50	840
7,50	0.70
	CTCAT
GCACC ACCGC GGACG GAGCA TGTGC GTTCT GGCCT CCGGC GCGTA CTGCG TTAAG	GAGTA
850 860 870 880 IB 890	900
TGTCG TGGCG ACAGT CCTCG TGATG GAGCT GACCG CCTAT TCCGT CCACC GCTGG	TATTA
ACAGC ACCGC TGTCA GGAGC ACTAC CTCGA CTGGC GGATA AGGCA GGTGG CGACC	ATAAT
910 920 930 940 950	960
GCACG GCCCC CTAGG CTGGG GCTGG CACAA GTCCC ATCAC GAAGA GCACG ACCAC	GCGTT
CGTGC CGGGG GATCC GACCC CGACC GTGTT CAGGG TAGTG CTTCT CGTGC TGGTG	CGCAA
970 980 990 1000 1010	1020
GGAGA AGAAC GACCT CTACG GCGTC GTCTT CGCGG TGCTG GCGAC GATCC TCTTC	ACCG1
CCTCT TCTTG CTGGA GATGC CGCAG CAGAA GCGCC ACGAC CGCTG CTAGG AGAAG	IGGCA
1030 1040 1050 1060 1070	1080
GGGCG CCTAT TGGTG GCCGG TGCTG TGGTG GATCG CCCTG GGCAT GACGG TCTAT	
CCCGC GGATA ACCAC CGGCC ACGAC ACCAC CTAGC GGGAC CCGTA CTGCC AGATA	CCCAA
ceede domin notine docte heade heade cined docine docine docine	
1090 1100 1110 1120 1130	1140
GATCT ATTIC ATCCT GCACG ACGGG CTTGT GCATC AACGC TGGCC GTTTC GGTAT	ATTCC
CTAGA TAAAG TAGGA CGTGC TGCCC GAACA CGTAG TTGCG ACCGG CAAAG CCATA	TAAGG
1150 1160 1170 1180 1190	1200
GCGGC GGGGC TATTT CCGCA GGCTC TACCA AGCTC ATCGC CTGCA CCACG CGGTC	GAGGG
CGCCG CCCCG ATAAA GGCGT CCGAG ATGGT TCGAG TAGCG GACGT GGTGC GCCAG	CTCCC
2050	1260
1210 1220 1230 1240 1250	
GCGGG ACCAC TGCGT CAGCT TCGGC TTCAT CTATG CCCCA CCCGT GGACA AGCTG CGCCC TGGTG ACGCA GTCGA AGCCG AAGTA GATAC GGGGT GGGCA CCTGT TCGAC	TTCGT
COCCC 16616 ACCCA GICGA AGCCG AAGIA GAIAC GGGGI GGGCA CCIGI 100A0	
1270 1280 1290 1300 1310	1320
GGATC TGAAG CGGTC GGGTG TCCTG CGCCC CCAGG ACGAG CGTCC GTCGT GATCT	
CCTAG ACTIC GCCAG CCCAC AGGAC GCGGG GGTCC TGCTC GCAGG CAGCA CTAGA	GACTA
TO	
1330 1340 1350 1360 1370	1380
CCCGG CGTGG CCGCA TGAAA TCCGA CGTGC TGCTG GCAGG GGCCG GCCTT GCCAA	CGGAC
	GCCTG
GGGCC GCACC GGCGT ACTTT AGGCT GCACG ACGAC CGTCC CCGGC CGGAA CGGTI	
GGGCC GCACC GGCGT ACTIT AGGCT GCACG ACGAC CGTCC CCGGC CGGAA CGGTT	
1390 1400 1410 1420 1430	1440
	1440 CCGTG

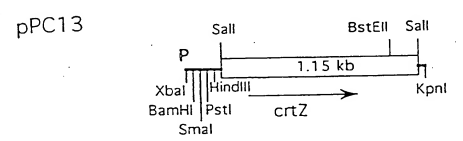
FIG. 17

	1450		1460		1470		1480		1490		1500
CGGCG	GGCGC	CTCGG	ACGGG	CATAC	TTGGT	CCTGC	CACGA	CACCG	ATTTG	GCGCC	
				GTATG							
	1510		1520		1530		1540		1550		1560
				ATCAG							
CCGAC	CTGGC	GGACT	TCGGC	TAGTC	CGCAC	CGCTG	ACCGG	GCTAG	TCCTC	CACGC	CAAGG
	1570		1580		1590		1600		1610		1620
CAGAC		GCGAA						CGACG		GGGCT	1620
CAGAC GTCTG	CATTC	GCGAA CGCTT	GGCTC	CGGGC GCCCG	CGGAT	ATGGC	TCGAT	CGACG GCTGC	GGCGG	GGGCT CCCGA	1620 GATGC
CAGAC GTCTG	CATTC	GCGAA CGCTT	GGCTC	CGGGC	CGGAT	ATGGC	TCGAT	CGACG GCTGC	GGCGG	GGGCT CCCGA	1620 GATGC
CAGAC GTCTG	CATTC	CGCTT	GGCTC	CGGGC	CGGAT	ATGGC	TCGAT	CGACG GCTGC	GGCGG	GGGCT CCCGA	1620 GATGC
GTCTG	CATTC	CGCTT	GGCTC	CGGGC	CGGAT	ATGGC	TCGAT	CGACG GCTGC	GGCGG	GGGCT CCCGA	1620 GATGC

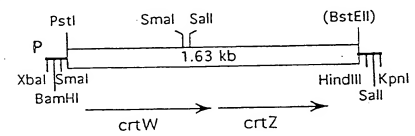
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pPC17



pPC17-3

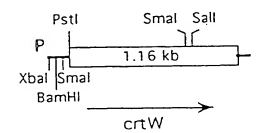
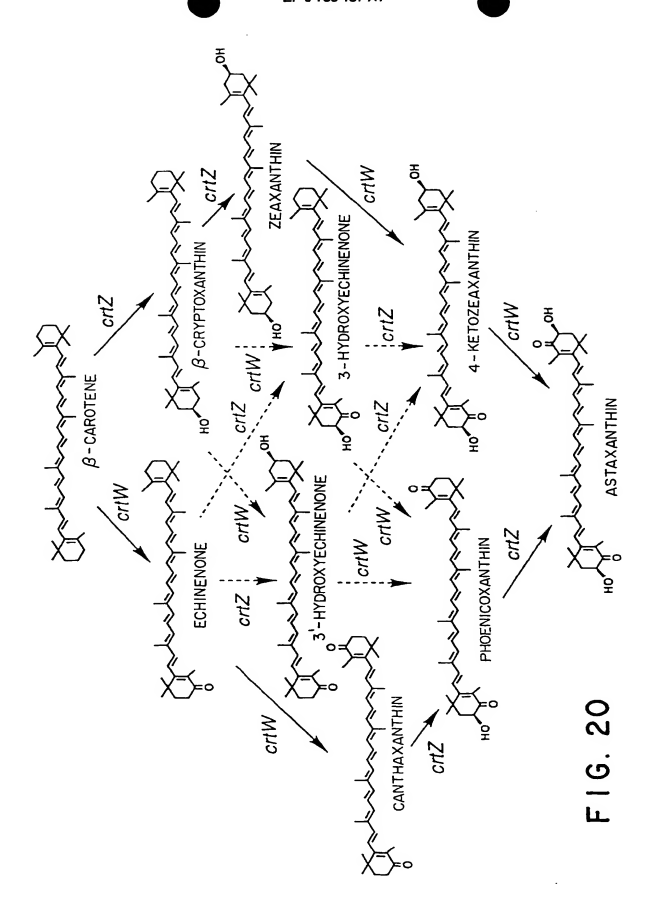


FIG. 19



INTERNATIONAL SEARCH REPORT

International application No.
PCT/JP94/02220

A. CLASSIFICATION OF SUBJECT MATTER Int. Cl ⁶ Cl2N15/00, Cl2P7/00									
	According to International Patent Classification (IPC) or to both national classification and IPC								
B. FIELDS SEARCHED									
Minimum documentation scarched (classification system followed by classification symbols)									
Int. C16 C12N15/00, C12P7/00									
Documentat	Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched								
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)									
CAS ONLINE, BIOSIS, WPI/WPIL									
C. DOCU	MENTS CONSIDERED TO BE RELEVANT								
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.						
E	WO, A, 9406918 (Gist-Broca	des NV.),	1-31						
	March 31, 1994 (31. 03. 94 &EP, A, 586751 & CA, A, 21								
	*	_							
A	EP, A, 474347 (Unilever Pl March 11, 1992 (11. 03. 92		1-31						
	& JP, A, 5-076347	•	•						
A	"Marine bacteria produced	astaxanthin"	1-31						
,	10th International symposi								
	abstract, CL11-3(1993)								
,									
	· ×								
	*								
Furthe	er documents are listed in the continuation of Box C.	See patent family annex.	·						
	categories of cited documents: an defining the general state of the art which is not considered	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand							
to be of	particular relevance locument but published on or after the international filing date	the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be							
"L" docume	L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other								
special	special reason (as specified) "Y" document of particular relevance; the claimed invention canno considered to involve an inventive step when the document referring to an oral disclosure, use, exhibition or other combined with one or more other such documents, such combinations.								
"P" docume	nt published prior to the international filing date but later than rity date claimed	being obvious to a person skilled in the art "&" document member of the same patent family							
	Date of the actual completion of the international search Date of mailing of the international search								
	ch 16, 1995 (16. 03. 95)	April 4, 1995 (04. 04. 95)							
Name and m	nailing address of the ISA/	Authorized officer							
Japa	Japanese Patent Office								
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